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SYNTHESIS AND STRUCTURE-PROPERTY CORRELATION OF PERFLUOROPOLYALKYLETHER BASE FLUIDS

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TECHNOLUBE PRODUCTS DIVISION LUBRICATING SPECIALTIES CO. 3365 E. SLAUSON AVENUE VERNON CA 90058



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13. ABSTRACT (Maximum 200 words)

Monomers for the preparation of the partially fluorinated model fluids were synthesized and limited investigations were carried out on their reactions. Direct fluorination processes were explored. A series of commercial and experimental perfluoropolyalkylether fluids was fully characterized and their thermal oxidative behavior in the presence of M-50 bearing steel alloy was determined. Computational treatments based on molecular mechanics, the input from the experimental data, and regression analysis led to the development of Quantitative Structure /Activity (Property) Relationships (QSAR). The developed expressions permit predictions, without synthesis of the candidate materials, of thermal oxidative stability, viscosity, viscosity/temperature and viscosity/molecular weight profiles.

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FOREWORD

This report was prepared by Technolube Products
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perfluoropolyalkylether fluids.

We also wish to acknowledge the contribution of Allied-Signal Inc., in the synthesis program under a subcontract.

1. EXECUTIVE SUMMARY

The goal of the subject contract was to develop structure/property (activity) relationships for perfluoropolyalkylether fluids to permit the identification of materials best suited for envisioned applications. The first portion of the program was performed at Ultrasystems Inc.; the majority of investigations were carried out at Technolube Products Division of Lubricating Specialties Co.

The goal of the synthesis effort was the preparation of partially fluorinated polyalkylether model fluids to be further fluorinated under subcontract with Allied-Signal Inc. The monomers, HO2CCF2OCF2CO2H, CloCCF2OCF2COCl, HOCH2CF2OCF2CH2OH and (COF)2 were successfully prepared, although the yields were lower than anticipated. Only limited effort was expended on the synthesis of partially fluorinated model fluids due to problems associated with direct fluorination and the sudden availability of a series of perfluoropolyalkylether fluids.

The major thrust of the work was centered on characterization of the fluids supplied by the Air Force, including determining their viscosity/temperature and viscosity/molecular weight profiles, and thermal oxidative stability behavior in the presence of metal alloys. The aim was to provide experimental data necessary for structure/activity relationships studies.

Based on the experimental input using computational chemistry in conjunction with multivariate regression analyses quantitative structure/activity (property) expressions were developed which allow the prediction of viscosity/temperature and viscosity/molecular weight profiles as well as the thermal oxidative stability of perfluoropolyalkylether fluids. The significance of this achievement is the ability to select the optimum compositions for specific applications without the need to synthesize a series of candidates. The validity of the approach was proven by the correct prediction of properties of known perfluoropolyalkylethers using the developed parameters and expressions.

2. INTRODUCTION

Perfluoropolyalkylethers are the only class of materials which possess thermal oxidative stability and associated viscosity temperature characteristics, such as to permit, at least in theory, operation between -70 and 400°C. It is clear from the literature data [Ref. 1-7] that in the perfluoropolyalkylether series of fluids the linear arrangements exhibit better viscosity/temperature profiles; i.e., have a higher viscosity index (VI) than the corresponding branched chain analogues. The most important factor in improving the low temperature properties and increasing the VI is a high oxygen/carbon (O/C) ratio. Yet it is the same factor which appears to lead to lowered thermal oxidative stability in the presence of metals [Ref. 8]. Inasmuch as these materials are being considered as high temperature turbine engine oils for the future generation of aircraft, this aspect is of utmost concern.

The current study was undertaken to develop valid premises for the selection of the best compromise fluids for specific application. The goal was to prepare a series of well characterized "model" fluids and to develop structure/property relationships to permit identification of optimum arrangements without actual synthesis, based on the experimentally determined viscosity/temperature profiles and thermal oxidative behavior in the presence of metals. The goal of the synthesis portion of the program was to concentrate on the preparation of partially

fluorinated intermediates. These were to be transformed by direct fluorination, (to be carried out by Allied-Signal, Inc. under a subcontract), into a series of model fluids. The perceived advantage of this approach, as compared to direct fluorination of multicomponent hydrocarbon precursors (pioneered by Exfluor Research Corp.) was the formation of a single component fluid with well defined end groups, as well as avoidance of rearrangements, bond scissions and related problems inherent to fluorination of hydrocarbons. Having fluorinated entities in a given molecule moderates the initial reaction and prevents the formation of by-products.

As work proceeded the low yields obtained of the required intermediates and the difficulties encountered in the preliminary attempts to form the actual partially fluorinated precursors of the model fluids, together with the disappointing results of the direct fluorination efforts on test materials by Allied-Signal, led to the abandonment of this approach. The deciding factor in taking this action was the availability of a series of experimental fluids from Exfluor Research Corp. which appeared to offer, based on ¹⁹F NMR and GC/MS studies a viable alternative to the representative perfluoropolyalkylether systems. Furthermore work proceeding at that time at Exfluor Research Corp., under the Air Force sponsorship, indicated clearly that "purer" experimental fluids with a larger variety of chain arrangements were forthcoming. Accordingly, the latter fluids were utilized in the investigations which provided the

experimental data for the development of quantitative structure/activity (property) relationships (QSAR) for predicting the viscosities and the thermal oxidative stabilities of perfluoropolyalkylether fluids. The developed expressions were based on molecular mechanics/molecular dynamics computations and multivariate linear regression analysis.

3. RESULTS AND DISCUSSION

The overall program was composed of a number of tasks which provide a logical breakdown for the orderly presentation of the data. Thus this portion of the report has been divided into: Synthesis and Related Aspects, Characterization and Study of the First Series of Experimental Perfluoropolyalkylether Fluids, Characterization and Thermal Oxidative Stability Testing of Chlorofluoropolyalkylether Fluids, Characterization and Study of the Second Series of Experimental Perfluoropolyalkylether Fluids, Development of Quantitative Structure/Viscosity Relationships, Thermal Oxidative Stability Investigations and Development of Quantitative Structure/Stability Relationships.

3.1 SYNTHESIS AND RELATED ASPECTS

The major synthesis effort under this contract was directed at the preparation of the monomers $O(CF_2CO_2H)_2$, $O(CF_2COC1)_2$, $O(CF_2CH_2OH)_2$ and $(COF)_2$. These monomers together with derivatives such as $O(CF_2COF)_2$, $O(CF_2I)_2$ and $O(CF_2CH_2Br)_2$, in conjunction with the commercially available hexafluoropropene oxide, potentially provide the starting materials for a series of different fluorinated and partially fluorinated perfluoropolyalkylethers or their precursors.

A small portion of the synthesis work was expended on the evaluation of the direct fluorination capability at Allied-

Signal. Originally the subcontract with Allied-Signal, Inc. was to provide the direct fluorination support and computational expertise.

3.1.1 Synthesis of Monomers

The key starting material for the model fluid synthesis was HO₂CCF₂CCF₂CO₂H. One of the approaches pursued was originally described by the group at Dow Corning [Ref. 9] and is described in Scheme I.

The major problem we have encountered with the process, given in Scheme I, was the low yield of the isolated first intermediate, Cl₂C=CClCCl=CClOC₂H₅, which ranged from 23 to 32%. The tedious fractionation required to obtain the pure product complicated matters further. The next two steps proceeded essentially in quantitative yields. The 2,2,5,5-tetrafluoro-3,4-dichlorofuran was obtained in a 52% yield. Its oxidation to the acid HO₂CCF₂OCF₂CO₂H was initially attempted in acetone, but yields were poor and isolation difficult. Using alkaline permanganate gave the desired product in a 61% yield. Although the starting material for the Scheme I was inexpensive, the overall yield for the reactions up to the furan was very low, approximately 11%.

SCHEME I

As mentioned earlier attempts to optimize the first step were largely unsuccessful. Treatment of dichloromaleic anhydride with SF4 afforded another route to 2,2,5,5-tetrafluoro-3,4-dichlorofuran [Ref. 10, 11]. The advantage here was the one step process. The drawback was the cost of the starting material and of SF4.

Unfortunately the desired furan was invariably admixed with the lactone O=C-CCl=CClCF2, at best in equal proportions. Combined yields of the two products amounted to approximately 70%.

Another treatment with SF4 was necessary to attain reasonable yields approximately 45-50%, of the furan. This compares to the 11-20% attained using the multi-step reaction. Some process optimization was attempted in both of the instances however, the main thrust was to determine the feasibility of the formation of the desired model fluids.

The synthesis of the alcohol HOCH₂CF₂OCF₂CH₂OH, took place smoothly, in a 81% yield from the diethyl ester, following the reported procedure [Ref. 13]. It is of interest that using the methyl ester incomplete reaction was achieved. The diethyl ester was obtained in a 75% yield from the free acid and in 94% yield from mixture of the corresponding diacyl chloride and anhydride. Attempts to prepare pure diacyl chloride, O(CF₂COCl)₂, invariably resulted in mixture of the diacyl chloride and anhydride.

Oxalyl fluoride, (COF)₂, was obtained in a 52% yield by treatment of oxalyl chloride with sodium fluoride following the procedure of Tullock and Coffman [Ref. 14]. Interaction of the diol HOCH₂CF₂OCF₂CH₂OH with SOBr₂ failed to give

BrCH₂CF₂OCF₂CH₂Br. The latter was one of the co-reactants required for the synthesis of certain classes of model fluids.

3.1.2 Synthesis of Model Fluid Precursors

The route to the intermediate hydroxy-terminated polyesters was outlined in the original proposal [Ref. 15], i.e.

$$\begin{array}{c|c} \text{O(CF}_2\text{CH}_2\text{OH)}_2 + \text{O[CF}_2\text{C(O)]}_2\text{O} \\ & & & & & & \\ \text{O} & & & & & \\ \text{O} & & & & & \\ \text{O} & & & & & \\ \text{HO[CH}_2\text{CF}_2\text{OCF}_2\text{CH}_2\text{OCCF}_2\text{CO]}_x\text{CH}_2\text{CF}_2\text{OCF}_2\text{CH}_2\text{OH} \\ \end{array}$$

The value of x is dependent on the stoichiometry of the two reagents. Initial reaction was carried out in refluxing tetrahydrofuran using a mixture of the anhydride and the corresponding acyl chloride with the diol $O(CF_2CH_2OH)_2$ (in the mole ratio of 1:1.5) in the presence of pyridine. Only unreacted diol and the acid $O(CF_2CO_2H)_2$ were observed after workup. A reaction performed in the absence of solvent in a sealed tube at $80^{\circ}C$ gave the expected oligomers.

A parallel approach to acid terminated polyesters i.e.

using a reaction of a mixture of anhydride and acyl chloride with the diol in a ratio of 7.5 to 1 at 85°C in the absence of solvent gave what appeared to be the expected product. However, in view of the lack of progress in the direct fluorinations carried by Allied-Signal, Inc., purification of neither of the above systems was carried out.

Based on work performed earlier at Ultrasystems [Ref 16], the reaction of hexafluoropropene oxide with oxalyl fluoride should proceed readily to give, under selected conditions, $FC(0) CF(CF_3) OCF_2 CF_2 OCF(CF_3) COF:$

The product distribution is strongly dependent upon the reaction conditions. Thus, at low temperatures it was assumed that the cesium/oxalyl fluoride complex should be the primary nucleophilic species which would then attack the secondary carbon atom of the oxide [Ref. 17]. Higher temperatures favor the oligmerization of hexafluoropropene oxide. These investigations, as evident from Table 1, show clearly that the use of low temperature does lead to promising yields of the desired product. Optimization of the process was not pursued in view of lack of progress in the direct fluorination and the sudden availability

TABLE 1

ATTEMPTED SYNTHESIS OF F(0)CCF(CF₃)[OCF₂CF(CF₃)]_xOCF₂CF₂O[CF(CF₃)CF₂O]_yCF(CF₃)COF

	Recovered HFPO, mmol	7.9	22.3	18.3
	Recc HFP(П
Products (GC area %) ^a	Desired ^b Product	22.6	24.1	7.6°
	(HFPO) ₄	6.5	6.6	8.4
roducts ((HFPO) ₃	31.5	59.1	8.4
Pr	(HFPO) ₂	22.4	1.6	25.6
ions	Time hr	18	18	7
Conditions	Temp °C	-23	-12	0 7 -78-RT 16
	HFPO mmol	18.9	84.3	183
tions	Time hr	77	2	2
Condi	Temp Time	-23	-12	0
	IG II		5.3	0.9
	CsF	9.45	2.63	7.83 6.0 0
	(COF) ₂ mmo1	9.45	21.3	39.9
	Run	н	2	2

a) Analyzed as methyl esters.

b) Desired product x + y = 0, i.e., $F(0)CCF(CF_3)OCF_2CF_2OCF(CF_3)COF$.

c) Major product (42.2%, analyzed by GC as methyl ester): $\frac{\text{MeCO}_2\text{CF}(\text{CF}_3)[\text{OCF}_2\text{CF}(\text{CF}_3)]_{\textbf{X}}\text{OCF}_2\text{CF}(\text{CF}_3)\text{CF}_2\text{O}[\text{CF}(\text{CF}_3)\text{CF}_2\text{O}]_{\textbf{y}}\text{CF}(\text{CF}_3)\text{CO}_2\text{Me; 29.5% (GC) x + y = 1; 9.2% x + y } }{3.5% \text{ x + y = 3.} }$

= 2;

Runs 3 and 4 were unsuccessful in leading to any appreciable copolymerization products.

of the desired experimental fluids from Exfluor Research Corporation.

3.1.3 Direct Fluorination Results

Two materials, tetraglyme and diglycolic acid, were subjected to direct fluorination by Allied-Signal, Inc. to determine the applicability of the process for the synthesis of the envisioned model fluids.

The 8-hour treatment of tetraglyme resulted in 10.7 g of product which after purification was still contaminated by hydrogen- and carbonyl-containing impurities. A corresponding sample prepared by Exfluor Research Corp. and obtained from NASA Lewis was free from these impurities.

The attempts to prepare O(CF₂COF)₂ from diglycolic acid were totally unsuccessful. The 0.95 g sample received, based on the analysis performed, did not contain the desired acid fluoride nor the corresponding lactone or anhydride. The presence of hydrogenated compounds was clearly shown. Based on the failure of the fluorination process, the low yield tedious synthesis of the precursors and most importantly the sudden availability of experimental fluids of different arrangements, the synthesis efforts were abandoned in favor of the utilization of the experimental fluids in the development of the structure/property relationships.

3.2 CHARACTERIZATION AND STUDY OF THE FIRST SERIES OF EXPERIMENTAL PERFLUOROPOLYALKYLETHER FLUIDS

Based on the results presented in Section 3.1 the feasibility of preparing model fluids at a reasonable cost in a reasonable time period became doubtful. At the same time the experimental fluids, prepared by Exfluor Research Corp. from hydrocarbon precursors, became available and consequently it was decided by the Project Engineer to determine whether these materials could be utilized in the development of meaningful structure/property relationships. These fluid compositions were mixtures of telomers having also different end-groups.

Furthermore, the hydrocarbon precursors for the experimental fluids, in the majority of instances, were not well characterized. In addition during the fluorination process a certain degree of rearrangement was to be expected. Thus, characterization of the fluids was of great importance.

3.2.1 Fluid Characterization

Four fluids: MLO 88-48, MLO 88-51, MLO 88-129, and MLO 88-132, prepared by direct fluorination by Exfluor Research Corp., were received from the Air Force. The originally assigned structures, together with the structures proposed, based on ¹⁹F NMR and GC/MS data, are given in Table 2. Also included in Table 2 are molecular weights (determined by osmometry and by ¹⁹F NMR end-group ratios), viscosity data (received from WL/MLBT), and summary of GC results (the retention times denoting the limits of the elution envelope). As noted earlier, each of the fluids

TABLE 2

ANALYTICAL DATA AND STRUCTURE CORRELATIONS FOR SELECTED PERFLUOROALKYLETHER COMPOUNDS³

Chemical Structure Assigned ^C
C ₃ F ₇ O(CF ₂ CF ₂ CF ₂ CF ₂ O) _n C ₃ F ₇ C ₄ F ₉ O(CF ₂ CF ₂ CF ₂ O) _n C ₃ F ₇ C ₂ F ₅ O(CF ₂ CF ₂ CF ₂ O) _n C ₃ F ₇
CF30(CF2CF201 _n CF3 C2F50(CF2CF201 _n CF3
CF301CF20CF2CF201nCF3 CF301CF20CF2CF201nCF20C2F5 plus others
CF ₃ O((CF ₂ CF ₂ O) ₃ CF ₂ O) _n CF ₃ plus others

These samples were received from AFML (WRDC/MLBT).

This was the structure given. 3

These structures were assigned based on ¹⁹F NMR and GC/MS anelysis. Û

d) This molecular weight was measured by osmometry. e) This molecular weight was calculated from $^{19}{\rm F}$ NMR based on CF $_3$ - and group and an internal segment. •

These data were provided by WRDC/MLBT.

These values pertain to GC elution envelope of the given fluid. Material turned white and failed to flow.

comprises a series of homologues, as illustrated by the actual GC traces given in Figures 1-4.

In the case of MLO 88-48, the specific end-groups assigned originally to the fluid appear to be absent, based on $^{19}\mathrm{F}$ NMR data. The GC trace (Figure 1) shows a repeating pattern of two low intensity peaks and one high intensity peak. It has been established from GC/MS that there are three types of homologues present: 1) $\mathrm{C_3F_7O}(\mathrm{CF_2CF_2CF_2CF_2O})_{\mathrm{n}}\mathrm{C_3F_7}$ (major component); 2) $\mathrm{C_4F_9O}(\mathrm{CF_2CF_2CF_2CF_2O})_{\mathrm{n}}\mathrm{C_3F_7}$;

3) $C_2F_5O(CF_2CF_2CF_2CF_2O)_nC_3F_7$. In Figure 1 the peak at 19.3 minutes was found to be due to a material terminated by C_3F_7 at both ends. At 19.7 minutes the termination is C_4F_9 ; at 20.3 minutes C_2F_5 and C_3F_7 . The listing of the revelant ions is presented in Table 3. The assignments for the specific components, which were differentiated by the end-groups, are based on the presence or absence of characteristic ions, for C_4F_9 , m/e 435; for C_3F_7 , m/e 385; and for C_2F_5 , m/e 335.

The 3100 number average molecular weight determined by osmometry is significantly lower than the 4300 value obtained from the $^{19}{\rm F}$ NMR. Small quantities of low molecular weight impurities could be responsible.

The experimental fluid, MLO 88-51, exhibited in its GC trace (Figure 2) the presence of two distinct compositions. Based on $^{19}{\rm F}$ NMR spectroscopy in conjunction with the GC/MS data, the major peak is CF30(CF2CF2O)nCF3 and the minor peak is

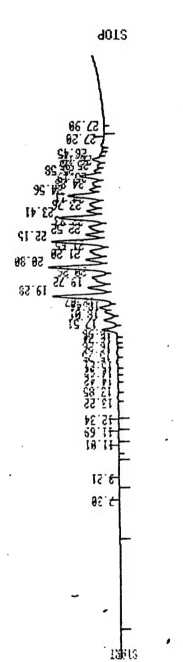
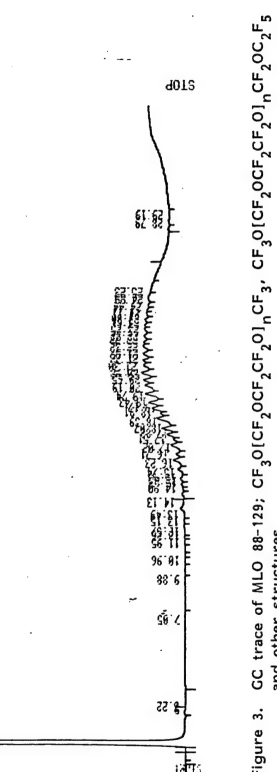


Figure 1. GC trace of MLO 88-48; $C_3F_7O[(CF_2)_4O]_nC_3F_7$, $C_4F_9O[(CF_2)_4O]_nC_3F_7$, $C_2F_5O[(CF_2)_4O]_nC_3F_7$

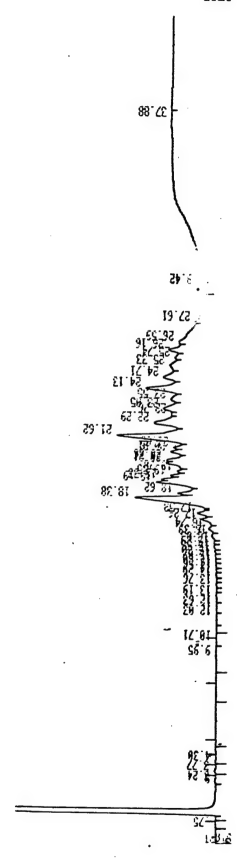


Figure 2. GC trace of MLO 88-51; CF₃O[CF₂CF₂O]_nCF₃, C₂F₅O[CF₂CF₂O]_nCF₃.





and other structures. Figure 3.



GC trace of MLO 88-132; CF₃O[(CF₂CF₂O)₃CF₂O]_nCF₃ and other structures.

TABLE 3

LISTING OF PROMINENT IONS SUPPORTING THE STRUCTURE ASSIGNMENT FOR MLO 88-48

Major Component: C₃F₇O[CF₂CF₂CF₂CF₂O]_nC₃F₇

Minor Components: $C_4F_9O[CF_2CF_2CF_2CF_2O]_nC_3F_7$

C₂F₅O[CF₂CF₂CF₂CF₂O]_nC₃F₇

<u>m/e</u>	<u>ion</u>
150	$CF_2CF_2CF_2$
169	C ₃ F ₇
197	CF2CF2CF2COF
219	C_4F_9
235	C ₃ F ₇ OCF ₂
335	$\mathtt{C_2F_5OCF_2CF_2CF_2CF_2}$
385	$\mathtt{C_3F_7OCF_2CF_2CF_2CF_2}$
435	$C_4F_9OCF_2CF_2CF_2CF_2$

 $C_2F_5O(CF_2CF_2O)_nCF_3$. The prominent MS peak assignments for the two compounds are given in Table 4.

The molecular weights as determined by osmometry and ^{19}F NMR are within experimental error. The molecular weight data, low temperature properties, and structural arrangements of MLO $^{88-48}$ and MLO $^{88-51}$ cannot be readily correlated. Ignoring the higher molecular weight of MLO $^{88-48}$ (due to the discrepancy in the two values), but based solely on the O to CF2 ratio, MLO $^{88-48}$ would be expected to exhibit a much higher viscosity at $^{-18}^{\circ}\text{C}$ than MLO $^{88-51}$. This would also be in line with the reported viscosities at 40 and $^{100}^{\circ}\text{C}$. On the other hand the well ordered structure of $^{-}(\text{CF}_2\text{CF}_2\text{O})_{\text{N}}$ — most likely promotes crystallization reflected by a drastic increase in viscosity at low temperature.

The GC data obtained for MLO 88-129, as evident from Figure 3, were poor. The $^{19}{\rm F}$ NMR spectroscopy indicated the presence of CF3 and C2F5 end-groups with the former predominating approximately in a 2:1 ratio. Thus, the major components seem to be CF30(CF2OCF2CF20)nCF3 and CF30(CF2OCF2CF20)nC2F5. The MS data support fully the $^{19}{\rm F}$ NMR assignments. The molecular weights, Table 2, determined by the two different methods, do not correspond closely. The only explanation for the higher value obtained by osmometry is chain branching.

MLO 88-132 could not be satisfactorily analyzed by GC/MS. The GC trace, Figure 4, points to many different arrangements; these could not be resolved. The ¹⁹F NMR data does not agree with the basic structure originally postulated. The

TABLE 4

LISTING OF PROMINENT IONS SUPPORTING THE STRUCTURE ASSIGNMENT FOR MLO 88-51

Major Component: CF₃O[CF₂CF₂O]_nCF₃

Minor Component: $C_2F_5O[CF_2CF_2O]_nCF_3$

<u>m/e</u>	<u>ion</u>
75	O-C-COF
135	CF ₃ OCF ₂
185	CF3CF2OCF2, CF3OCF2CF2
213	CF2CF2CCF2COF, CF2CCF2CF2COF
235	CF3CF2OCF2CF2
301	CF3OCF2CF2CF2
329	CF2CF2OCF2CF2COF
351	C ₂ F ₅ OCF ₂ CF ₂ OCF ₂ CF ₂
417	CF30[CF2CF20]2CF2CF2
445	[CF ₂ CF ₂ O] ₃ CF ₂ COF
467	C ₂ F ₅ [OCF ₂ CF ₂] ₃
561	[CF2CF2O]4CF2COF
677	[CF2CF2O]5CF2COF

arrangement $CF_3O[(CF_2CF_2O)_3CF_2O]_nCF_3$ is supported both by ^{19}F NMR and mass spectral data. Also there is a discrepancy in the molecular weights determined by ^{19}F NMR and osmometry. The lower value by NMR indicates branching. Based on the O:CF₂ ratio, the viscosities of MLO 88-129 should be lower than that of MLO 88-132, the experimental data verifies this assumption.

3.2.2 Stability Evaluations

To determine the effect of structural arrangement on the thermal oxidative stability, the perfluoropolyalkylether fluids MLO 88-48, -51, -129 and -132 described in Section 3.2.1, were subjected to thermal oxidative evaluations in the presence of M-50 steel bearing alloy. The results of these tests are summarized in Tables 5 and 6. As requested by the Project Engineer 15°C temperature increases starting with 260°C were employed to determine the stability as a function of temperature. Based on the results obtained, it can be seen that MLO 88-48, Rf0[CF2CF2CF2CF2O]nRf, undergoes steady degradation up to 330°C in direct agreement with Arrhenius rule of rate doubling for every 10°C. This would imply not the breakdown of the major component, but rather decomposition of an impurity.

The results for MLO 88-51, $R_fO[CF_2CF_2O]_nR_f$, indicate that between 300 and 315°C drastic increase in the degradation rate takes place. This break point for MLO 88-129, $R_fO[CF_2OCF_2CF_2O]_nR_f$, is between 275 and 290°C, whereas for MLO 88-132, $R_fO[(CF_2CF_2O)_4CF_2O]_nR_f$, the value lies between 300 and

TABLE 5

COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PERFLUOROALKYLETHER FLUIDS^a

Test No	Fluid Type	g	Temp °C	Volatiles mg/g	Coupon Weight 2	Reference
1	Krytox (MLO 71-6)	3.02	260	0.13	+0.1	2-90-101
2	Fomblin Z (Z25-P28)	3.17	260	57.4	-0.2	2-90-102
3	Demnum S20 (MLO 86-51)	3.74	260	1.0	+0.2	2-90-107
4	Brayco 814Z (MLO 78-80)	3.32	260	4.8	+0.2	2-90-108
5	$R_{f}^{O[(CF_{2})_{4}O]_{n}R_{f}}$ (MLO 88-48) $R_{f} = C_{2}F_{5}, C_{3}F_{7}, C_{4}F_{9}$	3.21	260	0.06	0.0	2-90-115
9	ditto	3.20	275	0.50	-0.1	5-90-8
13	ditto	3.16	290	1.55	+0.06	5-90-14
17	ditto	3.18	315	6.95	+0.3	5-90-60
24	ditto	3.19	330	21.7	+0.4	5-90-105
6	$R_{f}^{O(CF_{2}CF_{2}O)}_{n}R_{f}^{R}$ (MLO 88-51) $R_{f} = CF_{3}, C_{2}F_{5}$	3.22	260	0.50	0.0	2-90-116
10	ditto	3.22	275	0.52	0.0	5-90-9
14	ditto	3.22	290	0.53	+0.06	5-90-15
19	ditto	3.20	300	1.47	+0.3	5-90-96
18	ditto	3.20	315	63.7	0.0	5-90-61
21	ditto	3.20	315	157.6	-4.2	5-90-100
7	$R_f^{O[CF_2OCF_2CF_2O]}_{R_f} (MLO 88-129)$ $R_f = CF_3, CF_2OC_2F_5$	3.26	260	0.52	+0.1	2-90-134
11	ditto	3.24	275	2.1	+0.1	5-90-10
15	ditto	3.22	290	19.3	+0.4	5-90-24
8	$R_{f}^{O[(CF_{2}CF_{2}O)_{4}CF_{2}O]_{n}}R_{f}^{R}$ (MLO 88-132) $R_{f}^{O[(CF_{2}CF_{2}O)_{4}CF_{2}O]_{n}}R_{f}^{R}$	3.28	260	0.67	-0.1	2-90-135
12	ditto	3.23	275	1.7	+0.1	5-90-11
16	ditto	3.19	290	4.67	+0.5	5-90-25
20	ditto	3.22	300	9.63	+0.8	5-90-97
22	ditto	2.84	315	446.5	+0.4	5-90-101

⁽a) All the tests were performed in pure oxygen at the denoted temperature in the presence of an M-50 coupon over a 24 h period using the modified, scaled-down, sealed version of the AFML Micro-O-C-Test arrangement

TABLE 6

COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PFPAE FLUIDS IN THE PRESENCE OF M-50, $\rm O_2$ FOR 24 H.

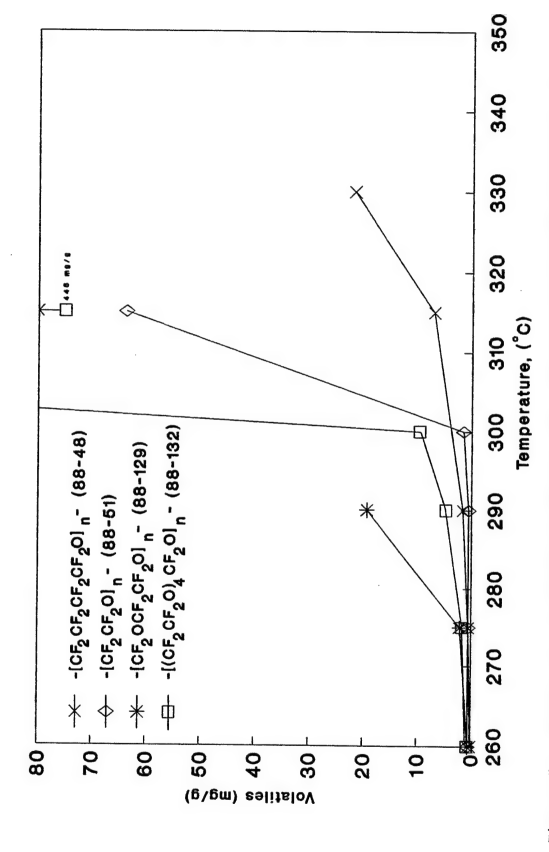
				Vola	Volatiles, mg/g	/مَ		
Fluid	MLO	260°C	275°C	290°C	3000	315°C	330°C	345°C
R_{f} o[$\mathrm{CF_{2}CF_{2}CF_{2}}$ o] $_{\mathrm{n}}$ R $_{\mathrm{f}}$	88-48	90.0	0.50	1.6		7.0	221.7	
$^{ m R_{ m f}o[CF_2CF_2o]_nR_{ m f}}$	88-51	0.50	0.52	0.53	1.5	63.7		
$^{ m R_fO[CF_2OCF_2CF_2O]_nR_f}$	88-129	0.52	2.1	19.3				
$^{ m R_fO[~(CF_2CF_2O)_4CF_2O]_nR_f}$	88-132	0.67	1.7	4.7	9.6	446.5		
Fomblin Z25-P28 $ m R_{f}O[CF_{2}O]_{m}[CF_{2}CF_{2}O]_{m}R_{f}$	1	57.4						
$\begin{array}{l} \text{Brayco 814Z} \\ \text{R}_{\text{f}}\text{O}[\text{CF}_{\text{2}}\text{O}]_{\text{n}}[\text{CF}_{\text{2}}\text{CF}_{\text{2}}\text{O}]_{\text{m}}\text{R}_{\text{f}} \end{array}$	78-80	4.8	62.2					
$ ext{Krytox}$ 143AC $ ext{R}_{ ext{f}}$ O $(ext{CF}(ext{CF}_3) ext{CF}_2 ext{O}]_{ ext{n}} ext{R}_{ ext{f}}$	71-6	0.13		4.0 ^a (288°C)	38.2 ^a (72 h)	44.0 47.5a (316°C)		
Krytox 143AC (pretreated)						0.2ª (316°C)		15.4 ^a (343°C)
Aflunox 2509 R _f O[CF(CF ₃)CF ₂ O] _n R _f						0.5 ^C (316°C)		
Demnum S20 $ m R_{f}O(CF_2CF_2CF_2O)_nR_f$	86-51 88-177	1.0 0.7d (68 h)			1.4		13.3	
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1								

(a) Data from AFML-TR-77-150. Test conducted for 72 h. (c) Data from NASA CR-180872. (d) Test conducted for 68 h.

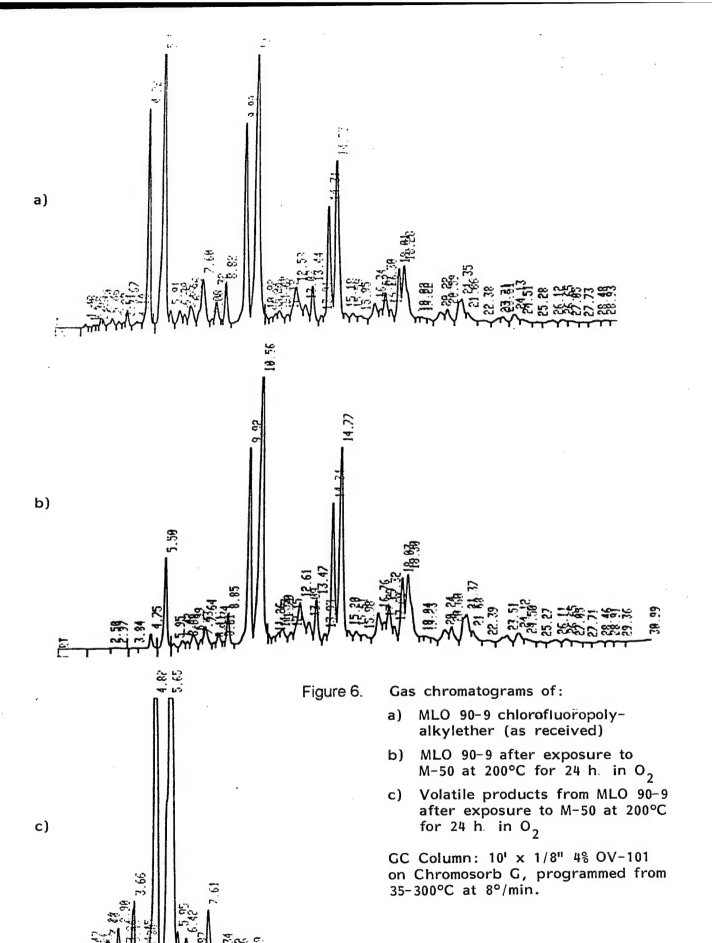
315°C. In view of the high degree of decomposition at 315°C, about 50%, actual degradation "onset" would appear to be closer to 300°C than 315°C. These data are presented graphically in Figure 5. The comparison given for commercial fluids at the top of Table 5 shows clearly the instability of the Fomblin Z family of fluids. The latter were the only materials tested having -OCF₂OCF₂O- segments believed to be responsible for the observed poor Fomblin Z stability [Ref. 8].

3.3 CHARACTERIZATION AND THERMAL OXIDATIVE STABILITY TESTING OF MLO 90-9 CHLOROFLUOROPOLYALKYLETHER FLUID

3.3.1 Characterization



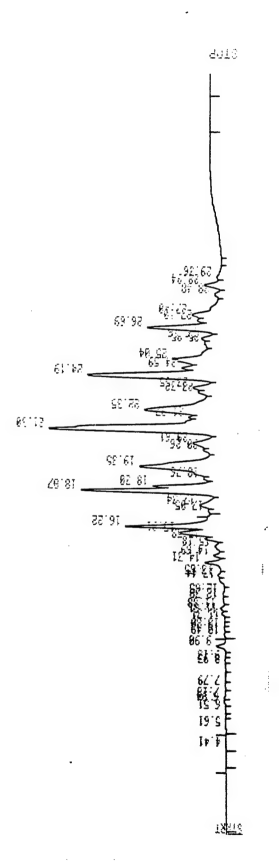
Comparison of thermal oxidative degradation data for MLO 88 series of experimental fluids. Figure 5.



For the other characterized chlorofluoropolyalkylether fluid, MLO 88-134, the postulated structure was $\text{ClCF}_2\text{CF}_2\text{O}[\text{CF}_2\text{CF}(\text{CF}_2\text{Cl})\text{O}]_n\text{CF}_3. \quad ^{19}\text{F NMR} \text{ analysis shows the end groups to be $-\text{CF}_2\text{CF}_2\text{Cl}$, $-\text{CF}_2\text{CF}_2\text{Cl}$, and $-\text{CF}_3$ in a ratio of $1.5:1.8:1. The gas chromatogram of the fluid is given in Figure 7. Based on GC/MS analysis, the main peak of the series consists of a mixture of <math>\text{ClCF}_2\text{CF}_2\text{O}[\text{CF}_2\text{CF}(\text{CF}_2\text{Cl})\text{O}]_n\text{CF}_2\text{CF}_2\text{CF}_2\text{Cl}}$ and $\text{ClCF}_2\text{CF}_2\text{O}[\text{CF}_2\text{CF}(\text{CF}_2\text{Cl})\text{O}]_{n+1}\text{CF}_3};$ the smaller peak appears to be $\text{ClCF}_2\text{CF}_2\text{O}[\text{CF}_2\text{CF}(\text{CF}_2\text{Cl})\text{O}]_n\text{CF}_2\text{CF}_2\text{CF}_2\text{Cl}}.$

3.3.2 Stability Evaluations

The fluid, MLO 90-9, was subjected to thermal oxidative stability testing at 200°C. As shown by the data listed in Table 7 at 200°C there was minimal or no degradation. Based on the GC analysis, the volatile products consisted mainly of the fluid's most volatile components, which apparently can be distilled out under vacuum at room temperature (compare Figures 6c and 6a). Consequently, the gas chromatogram of the fluid after exposure (Figure 6b) shows that most of the initially eluting components (up to ~6 minutes) have been removed. The fluid was retested at a higher temperatures 230°C and 260°C; the amount of volatiles collected increased slightly. The gas chromatogram of the vapor phase of the volatiles from the 230°C exposure (Figure 8a) shows some early eluting peaks (<2 minutes) not observed in the original fluid. Based on GC/MS analysis, two of the peaks have been identified as C1CF2CF=CF2 and C1CF2CF2CF2CF=CF2. The



GC column: 10' x 1/8" 4% OV-101 on Chromosorb G programmed from 35-300°C at $8^{\rm o}/\text{min.}$ Figure 7. Gas chromatogram of MLO 88-134 chloroperfluoropolyalkylether (as received).

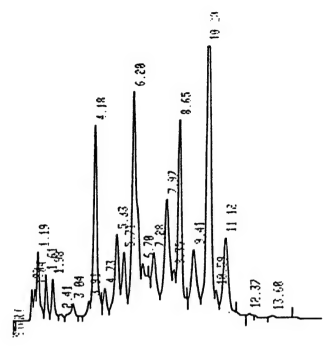
TABLE 7

THERMAL OXIDATIVE BEHAVIOR OF MLO 90-9
CHLOROFLUOROPOLYALKYLETHER FLUID^a

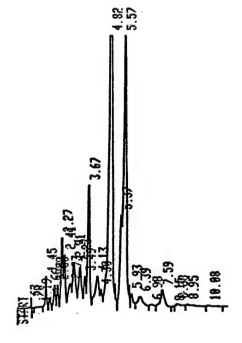
Test No	Wt., g	Temp °C	Volatiles mg/g	Coupon Weight Change, mg/cm ²	ID No.
23	3.22	200	13.1 ^b	+0.06	5-90-102
25	3.22	230	75.1 ^b	+0.01	5-90-106
26	3.23	260	183.3 ^b	+0.02	5-90-117

⁽a) All the tests were performed in pure oxygen at the denoted temperature in the presence of an M-50 coupon over a 24 hr period using the modified, scaled-down, sealed version of the AFML Micro-O-C-Test arrangement.

⁽b) Consisted mainly of the most volatile components of the original fluid.



a) Vapor phase, programmed from 35(2 mm hold)-300°C at 8°/min.



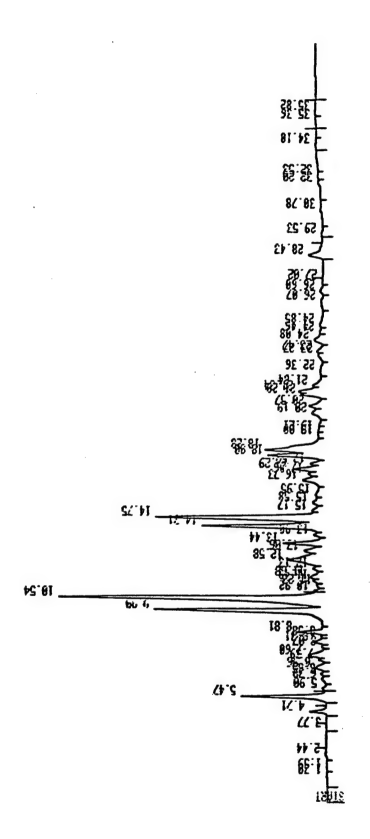
b) Liquid phase, programmed from 35-300°C at 8°/min.

Figure 8. Gas chromatograms of the vapor (a) and liquid (b) phases of the volatile products from MLO 90-9 after exposure to M-50 at 230°C for 24 h in O $_2$. GC column: 10' x 1/8" 4% OV-101 on Chromosorb G.

latter are definitely products of degradation; however, these compounds were produced only in trace quantities. The gas chromatogram of the liquid phase of the volatiles produced at 230°C (Figure 8) is very similar to that of the volatiles formed at 200°C (see Figure 6a). As mentioned earlier, these peaks are the volatile components of the original fluid. The GC trace of the fluid following the heat exposure at 230°C (Figure 9) shows depletion of the most volatile components.

3.4 CHARACTERIZATION AND STUDY OF THE SECOND SERIES OF EXPERIMENTAL PERFLUOROPOLYALKYLETHER FLUIDS AND COMMERCIAL FLUIDS

This series of fluids, listed in Table 8, was prepared using optimized direct fluorination procedures and more extensive post treatments than was the case for the materials discussed in Section 3.2. Inasmuch as these perfluoropolyalkylethers were to be employed in derivation of structure/property relationships the characterization and establishment of structures, on which to build the unavoidable assumptions, were mandatory. The commercial fluids listed in Table 9 provided both additional structural arrangements as well as different molecular weight fractions which permitted the development of molecular weight-viscosity curves. The latter were of great importance since none of the fluids are unimolecular and it was assumed that the use of an average molecular weight for a given spectrum of homologues does not invalidate computational treatments, and that the structural arrangements are the primary factors in imparting



GC column: 10' x 1/8" 4% OV-101 on Chromosorb G programmed from 35-300°C at $8^{\rm o}/{\rm min}$. Gas chromatogram of MLO 90-9 after exposure to M-50 at $230^{\circ}\mathrm{C}$ for 24 h in O_2 . Figure 9.

TABLE 8

PROPERTIES OF PERFLUOROPOLYALKYLETHERS

		Molec	Molecular Weighta	ighta	Viscosi	Viscosity ^b cSt/°C	၁ _{၀/}		
Fluid	MLO No.	Tech.	Exfl.	Osm.	-18	40	100	VIC	0/0
Fluorinated Pluronic L-31	91-87	2850	3000	2800	599	47.8	7.18	109	~2.3
Fluorinated Polyglycol P15-200	91-88	4900	5400	4250	480	54.1	8.67	137	~2.3
-(CF2CF2CF2OCF2O) $_{ m n}$ -	91-105	2700	2500	2500	1	20.8	5.04	183	2.5
- $(CF_2CF_2CF_2CF_2OCF_2O)_n$ -	91-106	2800	2600	2500	602	27.7	5.80	159	3.0
$-(CF_2CF_2CF_2O)_n^-$	91-126	3100	3000	2600	2370	52.0	7.73	114	4.0
Perfluoropoly(propylene oxide)	91-127	3250	3650	3100	7120	76.9	9.73	105	3.0
$-[\mathtt{CF}_2\mathtt{CF}_2\mathtt{OCF}_2\mathtt{CF}(\mathtt{CF}_2\mathtt{CF}_3)\mathtt{OCF}_2\mathtt{O}]_{\mathbf{n}}-$	91-131	2750	2900	2900	ı	22.6	5.00	155	2.3
$-\left\{ \mathtt{CF}_{2}\mathtt{CF}\left(\mathtt{CF}_{3}\right)\mathtt{OCF}_{2}\mathtt{CF}\left(\mathtt{CF}_{3}\right)\mathtt{OCF}_{2}\mathtt{O}\right\} _{n}-$	91-132	2700	3300	3050	11	28.4	00.9	165	2.3
-[$(CF_2CF_2O)_4CF_2O]_n$ -	91-157	1800	2100	2850	. 1	15.4	3.95	161	1.8
$-[\mathrm{CF_2CF}(\mathrm{CF_2OCF_2CF_2OCF_3})\mathrm{O]}\mathrm{n}-$	91-158	5650	4700	3950	4370	67.1	8.71	102	2.0
-(CF_2CF_2O) _n - (cross-linked)	91-159	1500	1300	3250	1190	31.1	5.53	115	2.0
- $(CF_2CF_2OCF_2O)_n$ -	91-160	2800	2300	2650	1	10.5	3.32	215	1.5
$-(CF_2CF_2O)_n$ (linear)	91-161	2450	2450	3000	р	16.7e	3.99e	537	2.0
- $[(c_2c_2c)_3c_F(c_3)o]_n$ -	91-162	1700	1750	2950	922	24.7	4.79	115	2.0

a) Number average molecular weights as calculated by Technolube and Exfluor from NMR data and as measured by vapor pressure osmometry by Technolube.

Viscosity determined by WL/MLBT except where noted.

Viscosity Index calculated by Technolube. After 24 and 72 h crystals formed in fluid, viscosity 4618 and 4714 cSt respectively These viscosity values were determined by Technolube. (a) (c) (c) (c) (c) (c) (d)

TABLE 9

PROPERTIES OF COMMERCIAL PERFLUOROPOLYALKYLETHERS

	\(\frac{1}{2}\)	Molecula	Molecular Weight	Viscos	Viscosity ^a cst/°C	sst/°c	144
Fluid	MLO NO.	NMK	OSM.	87	40	00T	\ \ \
Krytox 143AZ	66-92	1	1800	i	15.2	3.05	21
Krytox 143AA	66-91	1	2400	I	36.9	5.74	93
Krytox 143AB	42420p	4200	3200	ı	6.99	8.86	106
Krytox 143AB	66-93	I	3450	1	78.5	10.3	114
Krytox 143AC	53538b	6300	5250	ı	220	22.6	125
Krytox 143AC	71-6	0089	5400	33830	257	27.3	140
Krytox 143AD	66-95	ı	6650	I	420	40.9	148
Aflunox 606	88-295	1	2350	ł	28.6	4.84	84
Aflunox 1406	88-296	ı	2900	ı	53.2	7.64	107
Aflunox 2507	88-298	4300	3800	8550	95.2	12.0	117
Perfluoropoly(propylene oxide)	91-127	3250	3100	7120	6.97	9:73	105
Fluorinated Krytox 143AC	91-21	7250	0009	ı	329	33.3	143
Demnum S-20	ı	1	2500	ı	23.8	5.48	179
Demnum S-65	1	ı	4700	ı	63.8	12.6	201
Demnum S-100	1	ı	5700	1	86.1	16.4	206
Demnum S-200	1	I	7900	1	176	31.7	225

a) All viscosities listed were determined by Technolube except for the -18°C values which were determined by WL/MLBT.
b) Manufacturer batch number.

properties (i.e., viscosity, thermal oxidative stability) whereas the nature of the end-groups is of negligible influence.

3.4.1 Characterization

All the experimental fluids were subjected to detailed $^{19}{
m F}$ NMR analysis. The main objectives were to ascertain that the arrangements postulated were correct, to identify end-groups and their relative ratios, to determine molecular weights, and to identify the nature and proportion of impurities. The actual spectra are compiled in the Experimental Section. All the materials consisted of a series of analogues having different end-groups. This is clearly evident from the gas chromatograms presented in Figures 10-19 confirming the ¹⁹F NMR data. specific end-groups present in representative perfluoropolyalkylether analogues were identified by GC/MS in the first series of the experimental fluids. The data were discussed in Section 3.2 and will be not reiterated here. The commercial fluids likewise consist of a homologous series of differently end-capped telomers. This is illustrated by the chromatograms for Krytox family of materials given in Figures 20 and 21 (wherein the elution envelopes are associated with the average molecular weights). Corresponding chromatograms were obtained for representative Aflunox fluids as shown by Figures 22 and 23. The small adjacent peaks in Krytox fluids were in the past believed to be due to the presence of the hydrogen-terminated chains. Since Aflunox is hydrogen-free, as determined by ¹H NMR,



Figure 10. GC trace of MLO 91-105; $-(CF_2CF_2CF_2CF_2CF_2OCF_2O)_n$ -

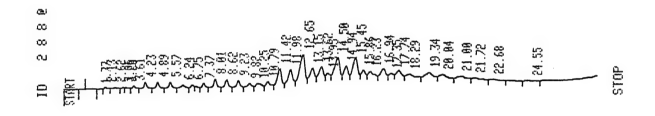


Figure 11. GC trace of MLO 91-106; $-(CF_2CF_2CF_2CF_2CF_2CF_2OCF_2O)_n$

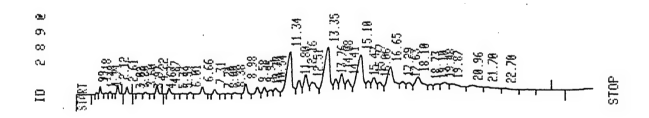


Figure 12. GC trace of MLO 91-126; $-(CF_2CF_2CF_2CF_2O)_n$ -



Figure 13. GC trace of MLO 91-127; $-[CF(CF_3)CF_2O]_n$ -

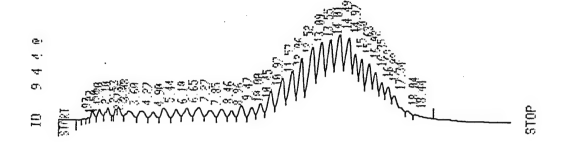


Figure 14. GC Trace of MLO 91-131; $-[CF_2CF_2CF_2CF_2CF_3]$ OCF_2O_n-1

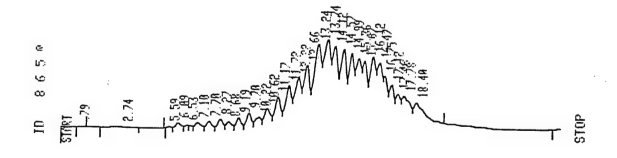


Figure 15. GC trace of MLO 91-132; $-[CF_2CF(CF_3)OCF_2CF(CF_3)OCF_2O]_n$ -

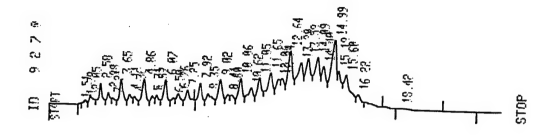


Figure 16. GC Trace of MLO 91-157; $-[(CF_2CF_2O)_4CF_2O]_n$ -

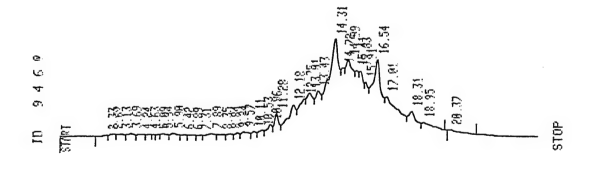


Figure 17. GC Trace of MLO 91-158; $-[CF_2CF(CF_2OCF_2CF_2OCF_2CF_3)O]_n$ -

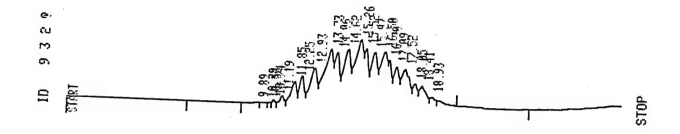


Figure 18. GC Trace of MLO 91-160; $-(CF_2CF_2OCF_2O)_n$ -

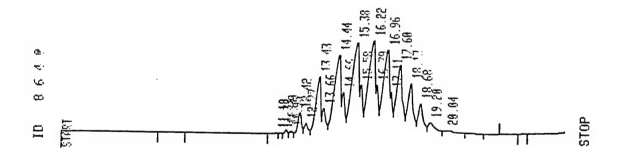


Figure 19. GC trace of MLO 91-161; $-(CF_2CF_2O)_n$ -

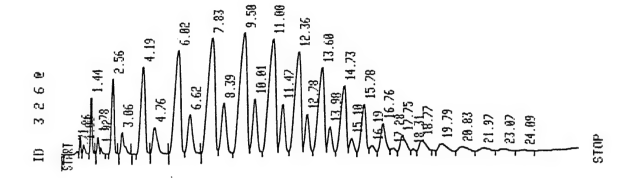


Figure 20. GC trace of Krytox 143AZ (ELO 66-92).

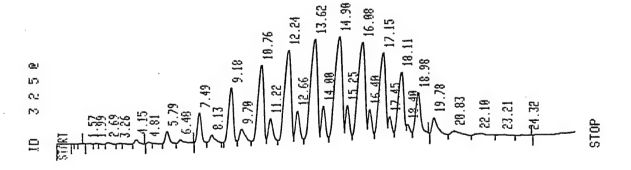


Figure 21. GC trace of Krytox 143AA (ELO 66-91).

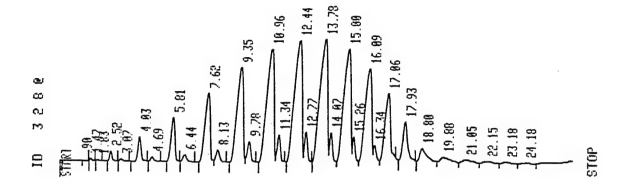


Figure 22. GC trace of Aflunox 606 (MLO 88-295).

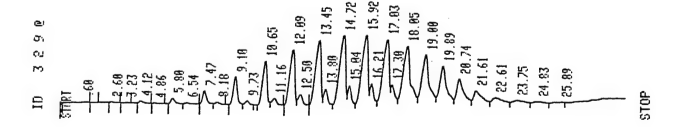


Figure 23. GC trace of Aflunox 1406 (MLO 88-296).

it is apparent that the observed pattern is due to different perfluoroalkyl end-groups. This aspect is clearly illustrated for Demnum S-20 in Figure 24 where each of the peaks in the repeating pattern was identified.

As mentioned earlier, all the experimental fluids

-Krytox 143AC, fluorinated Krytox 143AC, Demnum S-20, Demnum S
100 and Aflunox— were analyzed by NMR spectroscopy. These data

are given in the Experimental Section. It should be emphasized

that ¹H NMR spectra of Demnum, Aflunox and the fluorinated Krytox

did not show the presence of hydrogen. Hydrogen-containing end
group, -OCF(CF₃)H, was clearly evident in the ¹H NMR multi-scan

spectrum of Krytox 143AC.

Viscosity of any perfluoropolyalkylether is dependent both on its composition, i.e., structure, and its molecular weight. This number average molecular weight dependence is illustrated for the Krytox 143 series by the curve given in Figure 25. Regardless of the poly(perfluoropropene oxide) origin the molecular weight/viscosity profile is relatively consistent. Even the material obtained by direct fluorination of its hydrocarbon precursor was accommodated by the curve shown in Figure 26. Some discrepancy would be expected here since the structure, due to the synthesis path, is not as regular as that of the materials obtained by telomerization of hexafluoropropene oxide. As will be discussed later, it is the regularity which leads to a closely wound helix arrangement, which translates into the high viscosity characteristics.

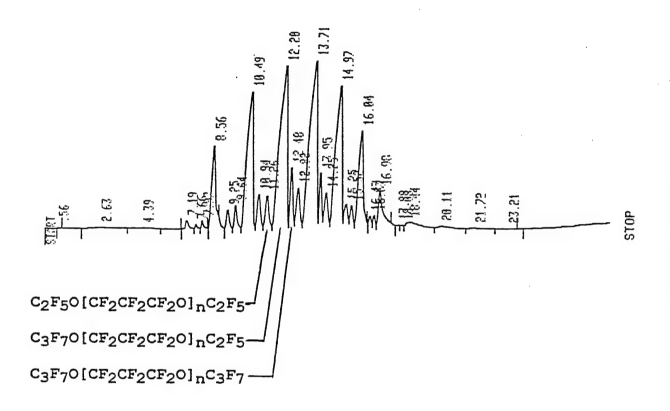


Figure 24. Gas chromatogram of fraction 1 from distillation of Demnum S-20 (MLO 88-177).

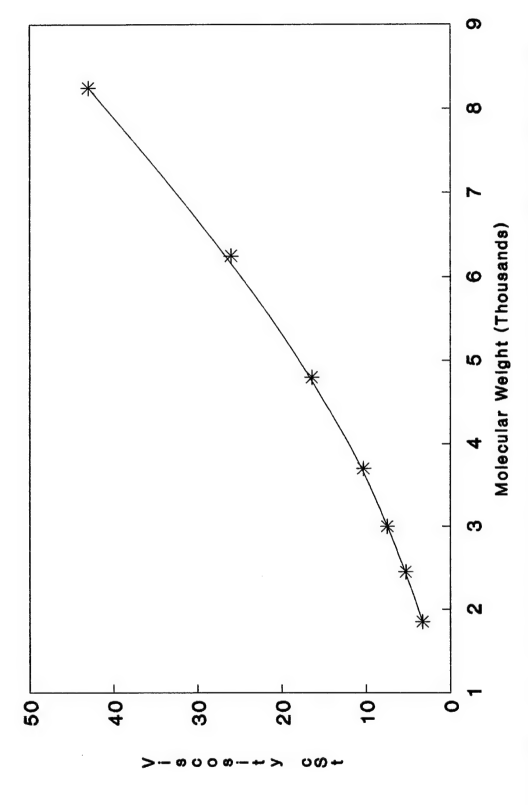
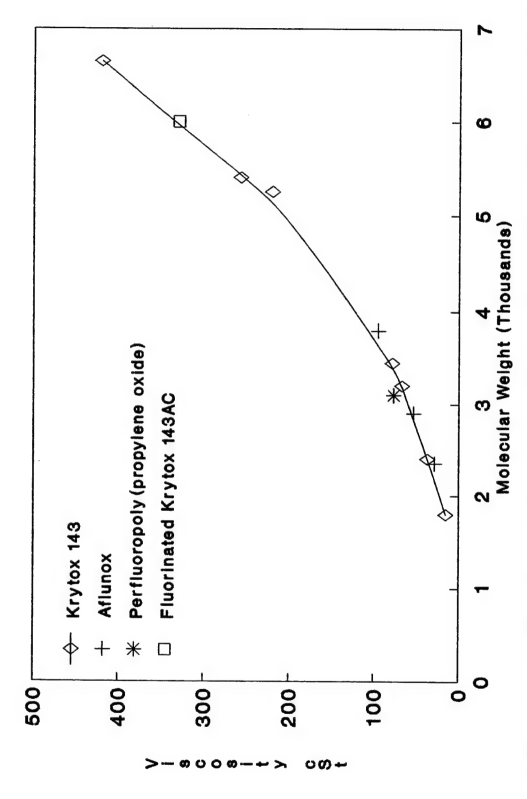


Figure 25. Plot of 100°C viscosity versus molecular weight for Krytox 143 fluids

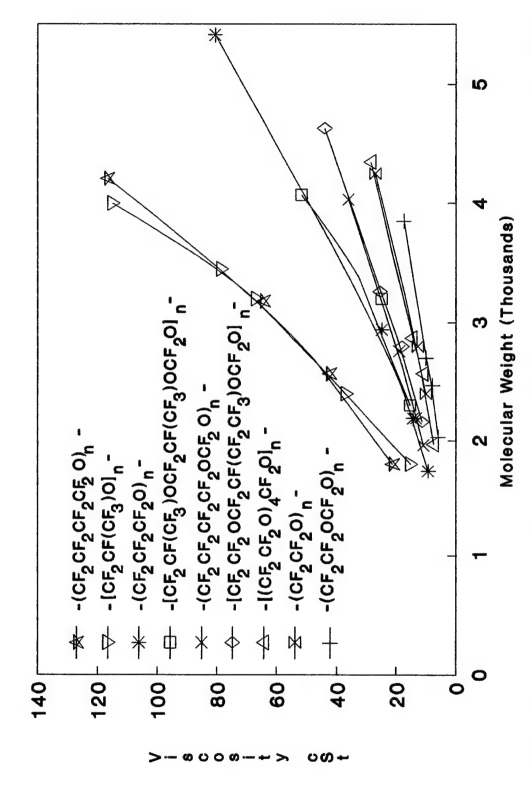


Plot of 40°C viscosity versus molecular weight for C3F70[CF(CF3)CF20]nC2F5 fluids from different sources. Figure 26.

To develop any viscosity/structure relationships requires the materials under consideration to be of identical molecular weights or molecular weight ranges. The obtaining of data for the poly(perfluoropropene oxide) family of fluids could be readily accomplished since a number of different molecular weight range fractions were available (see Table 9). In the case of Demnum and the experimental fluids, fractional distillation into more narrow molecular weight components was necessary to obtain the data points required for the viscosity/molecular weight plots. The distillation details are compiled in the Experimental Section (Table 60). The viscosity/molecular weight relationships for the fluids, employed in the development of the viscosity/structure relationships are given in Figures 27 and 28 for the 40°C and 100°C viscosities, respectively.

3.5 DEVELOPMENT OF QUANTITATIVE STRUCTURE/ACTIVITY (VISCOSITY) RELATIONSHIPS

None of the relatively high molecular weight perfluoropolyalkylethers that exhibit desirable viscosities and volatilities are unimolecular. All consist of telomer mixtures due to the nature of syntheses. An ability to correlate the structural arrangements and molecular weights, or rather molecular weight ranges, with viscosity characteristics would permit the identification of optimum perfluoropolyalkylethers for specific applications. Predictions of fluoroelastomer/chlorofluorocarbon swell [Ref. 18] and polysilahydrocarbon structure/viscosity [Ref. 19] were successfully accomplished



Viscosity/molecular weight profiles for perfluoropolyalkylethers at 40°C. Figure 27.

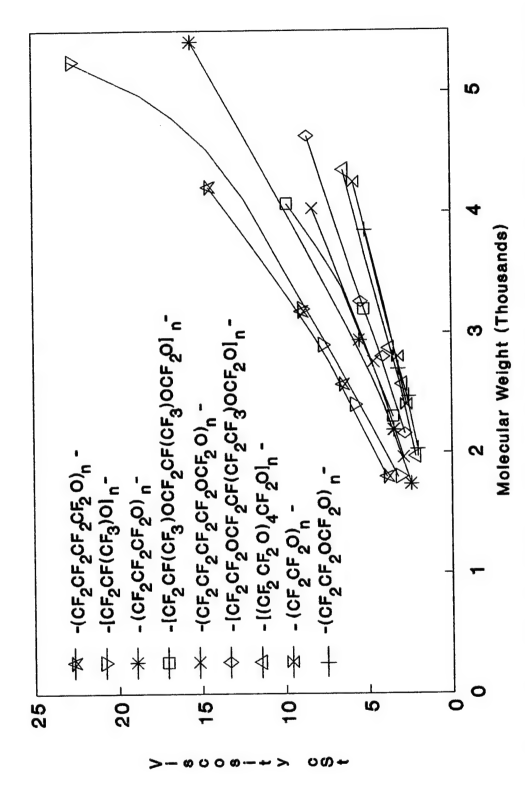


Figure 28. Viscosity/molecular weight profiles for perfluoropolyalkylethers at 100°C.

using a quantitative structure-activity relationships (QSAR). These studies provided the basis for the present undertaking.

For comparison of inherent viscosities and the determination of viscosity/structure relationships, as mentioned, materials of identical molecular weights or molecular weight ranges are required. The data used, compiled in Table 10, was obtained from plots given earlier in Figures 27 and 28. To develop QSAR expressions, the descriptors listed in Table 11 were employed in multivariate linear regression analysis. Using three descriptors with n = 9 (data points) gave r (Pearson correlation coefficient) of 0.963 and s (standard error estimate) of 4.27. The data point scatter is illustrated in Figure 29. With six descriptors r was raised to 0.985, s was 4.33; the actual plot is presented in Figure 30. In view of these results and the difficulty in encoding more complex structures, an approach involving energy parameters was investigated next.

The flexibility of a molecule, i.e., the ease to bend, is one of the major factors affecting the flow i.e., viscosity. In perfluoropolyalkylethers the carbon to oxygen ratio to a degree is related to this property. However, the latter relationship breaks down completely in the case of branched materials, as evident from the comparison of the relative viscosities of the isomers $-(CF_2CF_2CF_2O)_n$ and $-(CF(CF_3)CF_2O)_n$. In the case of longer CF_2 chains the C/O ratio provides an inadequate correlation as exemplified by the large difference in relative viscosities of $-(CF_2CF_2CF_2O)_n$ and $-(CF_2CF_2CF_2CF_2O)_n$.

TABLE 10
PERFLUOROPOLYALKYLETHER LISTING AND VISCOSITY DATA

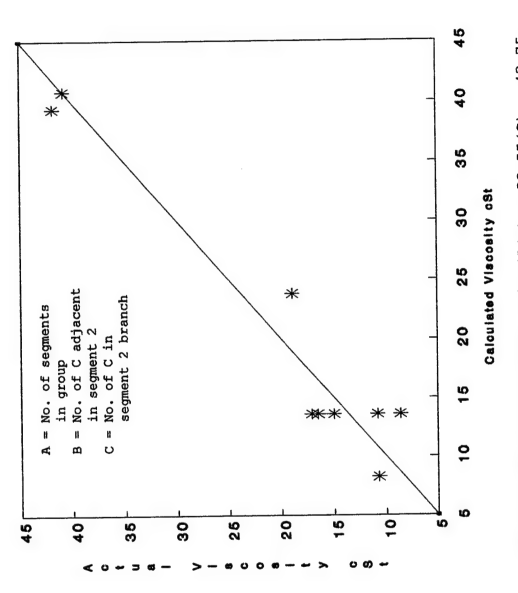
	MW 2500	200	MW 4000	000
Structure	Viscosity, cSt	V, cst	Viscosity, cSt	y, cst
	40°C	100°C	40°C	100°C
-(CF2CF20CF20)n-	8.61	2.78	18.5	5.5
$-(CF_2CF_2O)_n$	10.7	2.88	23.5	5.0
-[(CF_2CF_2O)4 CF_2O]n-	10.8	3.03	25.0	5.5
$-[\mathtt{CF}_2\mathtt{CF}_2\mathtt{CF}_2\mathtt{CF}_3)\mathtt{OCF}_2\mathtt{O}]_{\mathtt{n}}^-$	15.0	. 3,55	35.0	7.0
- (CF2CF2CF2OCF20) $_{ m n}$ -	16.5	4.05	35.7	8.15
- $[\mathtt{CF}_2\mathtt{CF}(\mathtt{CF}_3)\mathtt{OCF}_2\mathtt{CF}(\mathtt{CF}_3)\mathtt{OCF}_2\mathtt{O}]_n$ -	17.1	3.80	37.0	7.0
$-(CF_2CF_2CF_2O)_{n}$	18.9	4.30	49.0	0.6
$-[{\tt CF_2CF(CF_3)O]_n}^-$	40.8	6.05	115.0	12.0
$-(\text{CF}_2\text{CF}_2\text{CF}_2\text{O})_{n}$	41.8	6.36	106.0	13.0

TABLE 11

STRUCTURAL DESCRIPTORS CONSIDERED

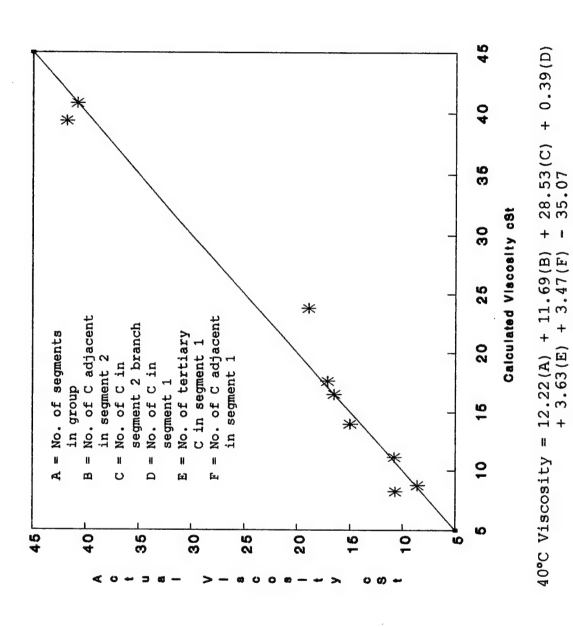
```
Carbon to oxygen ratio
Total number of carbons
Total number of oxygens
Total number of tertiary carbons
Number of segments in repeating group
Number of carbons in segment 1
Number of oxygens in segment 1
Carbon to oxygen ratio in segment 1
Number of carbons adjacent in segment 1
Number of tertiary carbons in segment 1
Number of carbons in branch of segment 1
Number of oxygens in branch of segment 1
Carbon to oxygen ratio in branch of segment 1
Number of carbons in segment 2
Number of oxygens in segment 2
Carbon to oxygen ratio in segment 2
Number of carbons adjacent in segment 2
Number of tertiary carbons in segment 2
Number of carbons in branch of segment 2
Number of oxygens in branch of segment 2
Carbon to oxygen ratio in branch of segment 2
```

Group and segment defined for e.g. $[(CF_2CF_2O)_4CF_2O]$ are as follows: Group = $[(CF_2CF_2O)_4CF_2O]$, Segment = $(CF_2CF_2O)_4$, CF_2O . The same values are assigned for type 2 as type 1 segments when only one type of segment is present.



 40° C Viscosity = 20.90(A) + 15.55(B) + 32.55(C) - 43.75

Calculated versus actual viscosity at 40°C as a function of three descriptors. Figure 29.



Calculated versus actual viscosity at 40°C as a function of six descriptors. Figure 30.

Molecular conformations of the perfluoropolyalkylethers presented in Figures 31-34 illustrate these aspects. Of special interest are the closely packed helical structures exhibited by the branched arrangements namely:

CF30[CF2CF2OCF2CF(CF2CF3)OCF20]6CF3,

CF₃O[CF₂CF(CF₃)OCF₂CF(CF₃)OCF₂O]₆CF₃ and CF₃O[CF₂CF(CF₃)O]₁₄CF₃. Another oligomer of the last material was reported by Pacansky et al [Ref. 20]. The coiling effect is absent in the short chain analogues [Ref. 21]; this aspect illustrates the inherent danger in developing theories regarding the behavior of polymers based on low molecular weight model compounds.

It is obvious that bonding energies, in particular bending energies, are major factors contributing to viscosity characteristics of a given material. To calculate the energy parameters, a number of assumptions and simplifications had to be made. Thus, it was assumed that the behavior of a unimolecular fluid parallels that of a telomer mixture of the same number average molecular weight. Furthermore, that the end-groups as such do not affect viscosity. Using these constraints the bending energies were calculated for the 2500 and 4000 molecular weight systems. The molecular weights of actual arrangements were selected as close as possible to these limits. These data are listed in Table 12.

Using C/O ratios and bending energies the expressions given below were developed for viscosities of the 2500 and 4000 molecular weight series at 40 and 100°C.

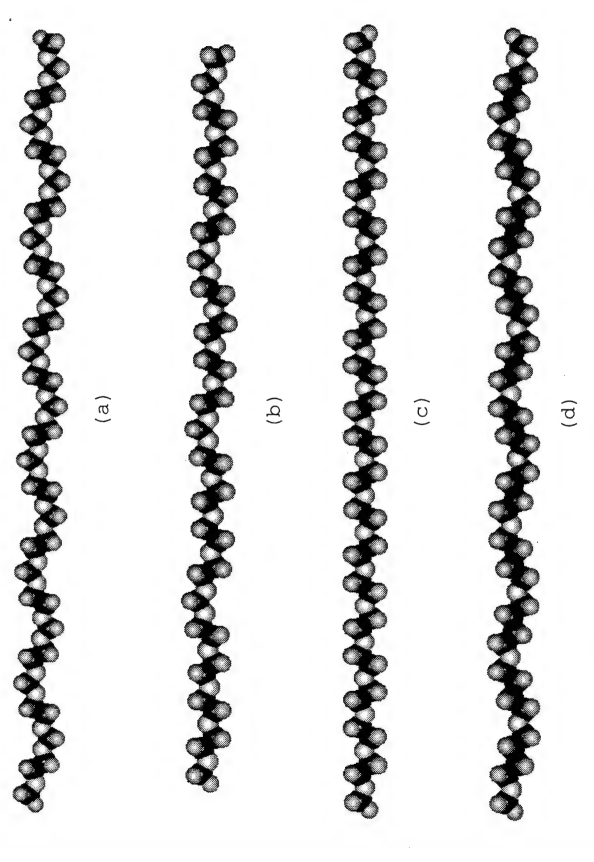


Figure 31. The conformations obtained with molecular mechanics for perfluoropolyalkylethers:
(a) CF3O(CF2CF2O)13CF3, (b) CF3O(CF2CF2O)4CF3,
(c) CF3O(CF2CF2O)20CF3, (d) CF3O(CF2CF2CF2O)11CF3.

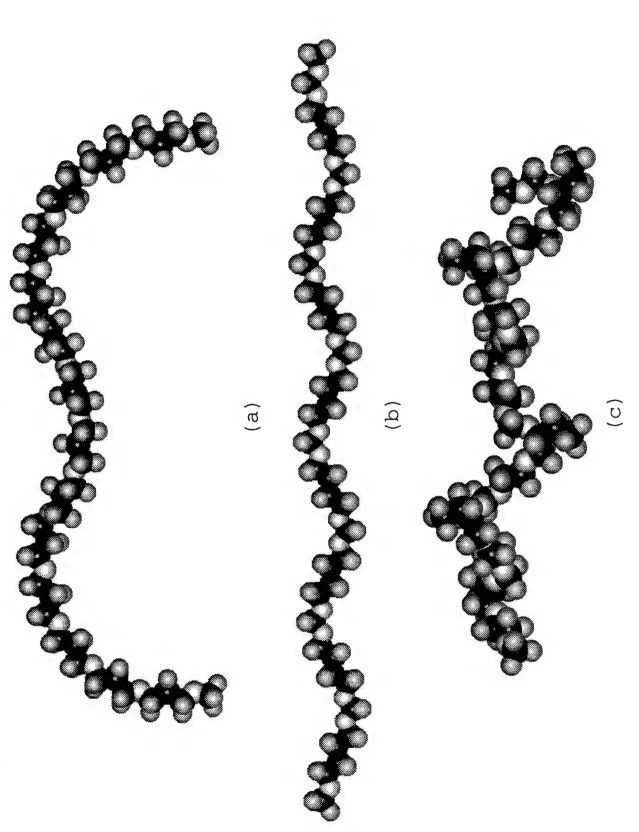
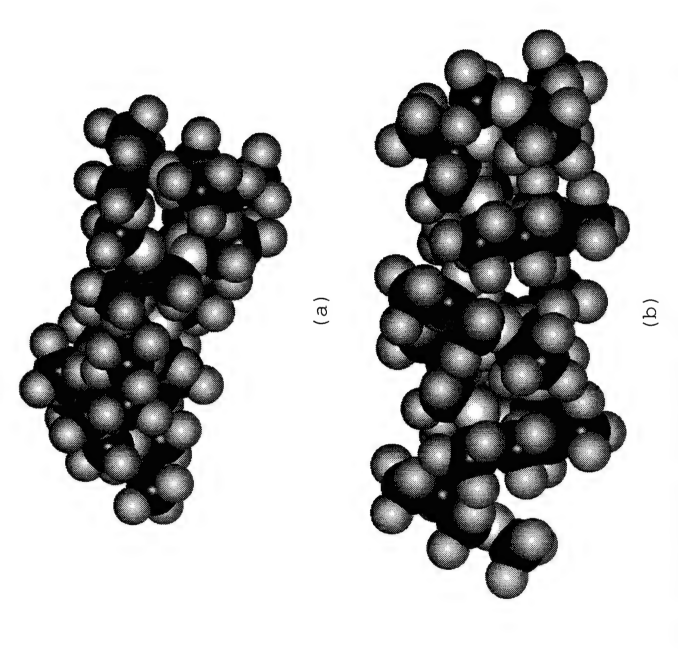


Figure 32. The conformations obtained with molecular mechanics for perfluoropolyalkylethers: (a) CF3O(CF2CF2CF2CF2CF2CF2OCF2O)8CF3, (b) CF3O(CF2CF2CF2CF2OCF2O)8CF3, (c) CF3O(CF2CF2CF2CF3)0CF2O]6CF3



The conformations obtained with molecular mechanics for perfluoropolyalkylethers: (a) CF30[CF2CF(CF3)0]14CF3. Figure 33.

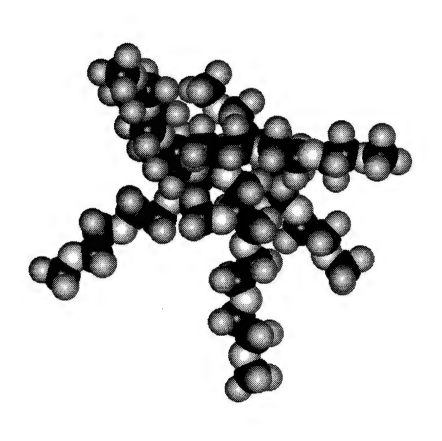


Figure 34. The conformation obtained with molecular mechanics for perfluoropolyalkylether: $CF_{30}(CF_{20}CF_{20}CF_{20}CF_{30}CF_{30})$

TABLE 12

PERFLUOROPOLYALKYLETHER BENDING ENERGY DATA

Structure		MW ~2500	500		MW ~4000	0
	п	MM	BEND (kcal mol-1)	ď) MM	BEND (kcal mol-1)
CF30 (CF2CF20CF20) nCF3	13	2520	13.4	21	3977	21.1
CF30 (CF2CF20) nCF3	20	2474	11.3	33	3983	18.1
$\mathtt{CF}_3\mathtt{O}$ [(CF2CF2O) $\mathtt{4}\mathtt{CF}_2\mathtt{O}$] $\mathtt{n}\mathtt{CF}_3$	4	2274	11.9	7	3865	20.5
${\tt CF}_3{\tt O}({\tt CF}_2{\tt CF}_2{\tt CF}_2{\tt CF}_3){\tt OCF}_2{\tt O}]_{\tt n}{\tt CF}_3$	9	2542	17.0	10	4135	28.3
$ ext{CF}_3 ext{O}$ (CF2CF2CF2OCF2O) $_{ ext{n}}$	ω	2410	15.6	14	4103	26.9
$\mathtt{CF}_3\mathtt{O}$ [$\mathtt{CF}_2\mathtt{CF}$ (\mathtt{CF}_3) $\mathtt{OCF}_2\mathtt{O}$] \mathtt{nCF}_3	9	2542	18.9	10	4135	30.3
$ ext{CF}_3 ext{O}\left(ext{CF}_2 ext{CF}_2 ext{O} ight)_{ ext{D}} ext{CF}_3$	14	2478	13.6	23	3973	22.4
$\mathtt{CF}_3\mathtt{O}\left[\mathtt{CF}_2\mathtt{CF}\left(\mathtt{CF}_3\right)\mathtt{O}\right]_{\mathtt{n}\mathtt{CF}_3}$	14	2478	25.9	23	3973	48.3
$\mathtt{CF}_3\mathtt{O}$ (CF2CF2CF2O) \mathtt{n} CF3	11	2530	15.7	18	4043	25.1

log 40°C Viscosity (MW,2500) = 0.244(C/0) + 0.022(BEND) + 0.279r = 0.992, s = 0.04, n = 9

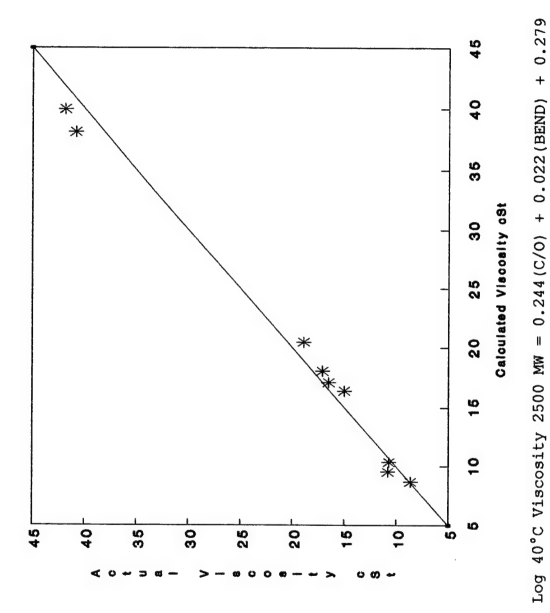
log 100°C Viscosity (MW,2500) = 0.137(C/O) + 0.011(BEND) + 0.084r = 0.990, s = 0.02, n = 9

log 40°C Viscosity (MW, 4000) = 0.278(C/O) + 0.013(BEND) + 0.572r = 0.991, s = 0.04, n = 9

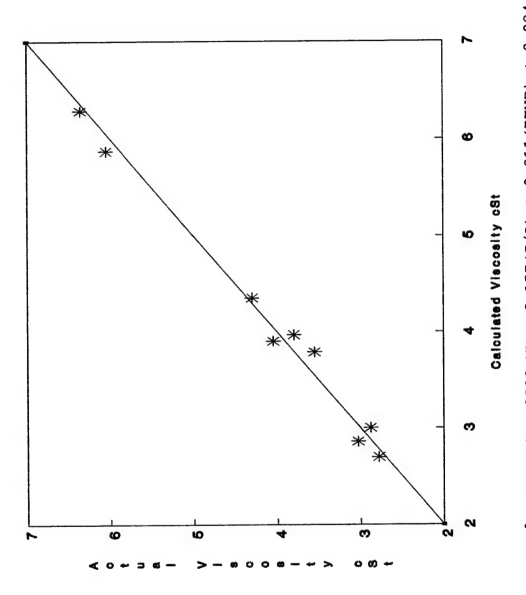
log 100°C Viscosity (MW, 4000) = 0.158(C/O) + 0.006(BEND) + 0.340r = 0.979, s = 0.04, n = 9

The corresponding graphical presentations are depicted in Figures 35-38. As evident from the high r values using just the two parameters, gives an unexpectedly good correlation in particular considering the assumptions made in developing these expressions. The actual graphs also show relatively low scatter.

To test the validity of the above relationships, analogous expressions were developed using eight of the materials. The viscosity of the excluded perfluoropolyalkylether was then calculated from these equations. This was done for each member of the series. The comparisons between the actual (obtained from the experimental plots given in Figures 27 and 28) and calculated viscosities are compiled in Table 13. The agreements, with the exception of $-(CF_2CF(CF_3)O)_n$ -, are reasonably close. It is believed, based on the conformation presented in Figure 33, that this perfluoropolyalkylether due to its tightly coiled structure, deviates strongly from the other

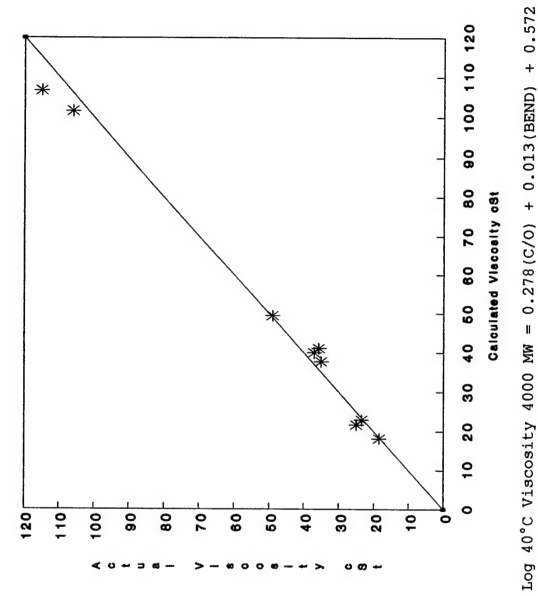


Calculated versus actual viscosity at 40°C of ~2500 MW perfluoropolyalkylethers as a function of the carbon to oxygen ratio and the bending energy. Figure 35.



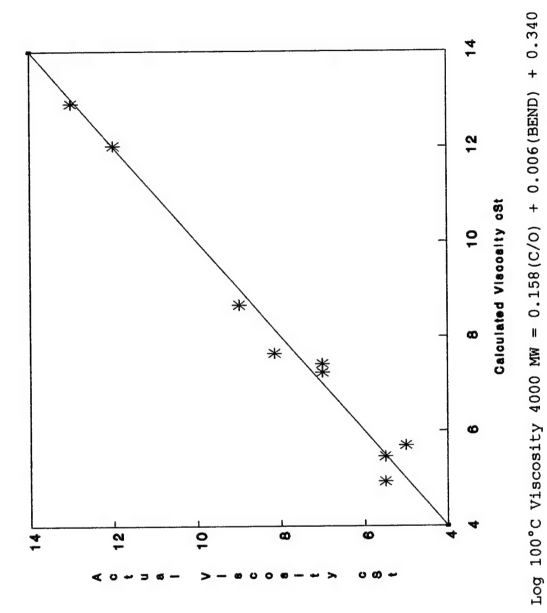
Log 100°C Viscosity 2500 MW = 0.137(C/O) + 0.011(BEND) + 0.084

Calculated versus actual viscosity at $100^{\circ}\mathrm{C}$ of $\sim 2500~\mathrm{MW}$ perfluoropolyalkylethers as a function of the carbon to oxygen ratio and the bending energy. Figure 36.



Calculated versus actual viscosity at 40°C of ~4000 MW perfluoropolyalkylethers Figure 37.

as a function of the carbon to oxygen ratio and the bending energy.



Calculated versus actual viscosity at 100°C of ~4000 MW perfluoropolyalkylethers as a function of the carbon to oxygen ratio and the bending energy. Figure 38.

TABLE 13

VISCOSITIES CALCULATED FROM EQUATIONS DERIVED FROM DATA EXCLUDING THE FLUID OF INTEREST

	MW 2	500 Visc	2500 Viscosity, cSt	St	MM	4000 Vi	4000 Viscosity, cSt	cst
Structure	4(40°C	100°C	၁	40°C	၁ွ	10	100°C
	actual	calc	actual	calc	actual	calc	actual	calc
- (CF2CF20CF20) n-	8.61	8.75	2.78	2.66	18.5	18.1	5.5	4.7
$-(CF_2CF_2O)_n^-$	10.7	10.3	2.88	3.05	23.5	22.8	5.0	5.9
-[(CF2CF20) $_4$ CF20] $_{\mathrm{n}}$ -	10.8	9,15	3.03	2.80	25.0	20.8	5.5	5.4
- $[CF_2CF_2OCF_2CF (CF_2CF_3)OCF_2O]_n$ -	15.0	16.6	3.55	3.83	35.0	38.0	7.0	7.3
$-(CF_2CF_2CF_2OCF_2O)_n$	16.5	17.2	4.05	3.89	35.5	41.8	8.0	7.6
- $[\mathrm{CF}_2\mathrm{CF}(\mathrm{CF}_3)\mathrm{OCF}_2\mathrm{CF}(\mathrm{CF}_3)\mathrm{OCF}_2\mathrm{O}]_\mathrm{n}$ -	17.1	18.3	3.80	4.02	37.0	40.5	7.0	7.5
$-(CF_2CF_2CF_2O)_n$ -	18.9	21.0	4.30	4.37	49.0	49.6	0.6	9.8
$-\left\{\text{CF}_{2}\text{CF}\left(\text{CF}_{3}\right)\text{O}\right\}_{n}$	40.8	31.4	6.05	5.39	115.0	73.6	12.0	12.3
- (CF2CF2CF2O) $_{ m n}$ -	41.8	35.8	6.36	6.10	106.0	7.06	13.0	12.6

arrangements. Consequently, an equation derived without an input from a related conformation would not be expected to provide valid results. Allowing for this inconsistency the relationships thus developed, using as the only parameters carbon to oxygen ratios and the bending energies, provide surprisingly reliable expressions for predicting viscosity/temperature and viscosity/molecular weight profiles. This capability is of great value in assessing the behavior of potential lubricants without a need to synthesize the actual materials. These techniques permit also accurate predictions for an increase in viscosity with increase in molecular weight for known compositions. It is apparent from the molecular conformations and related considerations that any predictions derived by extrapolation from low molecular weight arrangements, i.e. model compounds are unreliable if not meaningless.

3.6 THERMAL OXIDATIVE STABILITY INVESTIGATIONS OF FULLY CHARACTERIZED PERFLUOROPOLYALKYLETHER FLUIDS

This program, as mentioned in preceding sections, involved four commercial fluids, Krytox 143AC, Aflunox, Demnum S-20 and Demnum S-100, as well as a series of perfluoropolyalkylether experimental fluids. None of these materials are unimolecular; all consist of telomer mixtures due to the nature of the synthesis. The end groups tend to vary. These aspects were discussed earlier in this report. It is believed that insofar as stability is concerned the nature of the end groups whether CF3O, C2F5O, C3F7O, etc., is of a minor

importance compared to the effect of the type of repeating segments. On the other hand, impurities do affect the stability. This is obvious in comparing the thermal oxidative behavior of the first series of experimental fluids, listed in Table 6, with the data for the MLO 91 series discussed below and listed in Table 14. It must be stressed that perfluoropolyalkylethers exhibit very high thermal oxidative stabilities in the absence of metals i.e., when exposed to high temperatures in oxygen in glass. However, metals promote greatly the fluid degradation [Ref. 8] and lower significantly the degradation onset. of the envisioned application, namely a contact of the fluid at elevated temperatures with components manufactured from M-50 alloy or a related bearing material, all the evaluations were carried out in the presence of this metal. To facilitate the discussion the tabulation of experimental details for each test is given in the Experimental Section.

From the data presented in Figure 39 it is apparent that the commercial fluids, Krytox and Demnum, contain impurities which lower the degradation onset compared to materials given stabilization treatment. The Aflunox series, although believed to be prepared by the same or related process to Krytox, exhibit much better stability. It has been established that in Krytox fluids the impurities responsible for the decreased stability consist of hydrogen-terminated chains -OCF(CF3)H [Ref. 8]. The nature of the impurities in Demnum is unknown, although in view of the synthesis process [Ref. 5] residual hydrogen is again

TABLE 14

FLUIDS STUDIED

Degradation ^a	Temp°C	<288	325	<300	315	280	285	300	310	310	310	310	315
Molec weight	Osm.	5400	0009	2500		2850	2650	3050	3000	3950	2500	2500	2600
Molec	NMR	6800	7250	2750		1800	2800	2700	2450	5650	2700	2800	3100
	Origin	Krytox 143ACb	F-Krytox 143ACC	Demnum S-20d	T-Demnum S-20e	-f	1 I	J.	; 4-1	¥ 1	÷.	¥ 1	-f
	Fluid	$-[CF_2CF(CF_3)O]_n$	$-[CF_2CF(CF_3)O]_{\mathbf{n}}$	$-(CF_2CF_2O)_n$	$-(CF_2CF_2CF_2O)_n$	-[(CF_2CF_2O) $_4CF_2O$] $_n$ -	$-(CF_2CF_2OCF_2O)_n^-$	$-$ [CF2CF (CF3) OCF2CF (CF3) OCF2O $]_{ m n}-$	- (CF2CF20) n =	- [CF2CF (CF2OCF2CF2OCF2CF3) 0] n-	$-(CF_2CF_2CF_2OCF_2O)_n$	$-(CF_2CF_2CF_2CF_2OCF_2O)_n$	$-(CF_2CF_2CF_2O)_n^-$
	ID	71-6	91-21	88-177	88-177T	91-157	91-160	91-132	91 - 161	91-158	91-105	91-106	91-126

This is the degradation onset temperature at which volatile condensables amounted to $0.5 \pm 0.25 \text{ mg/g}$. a)

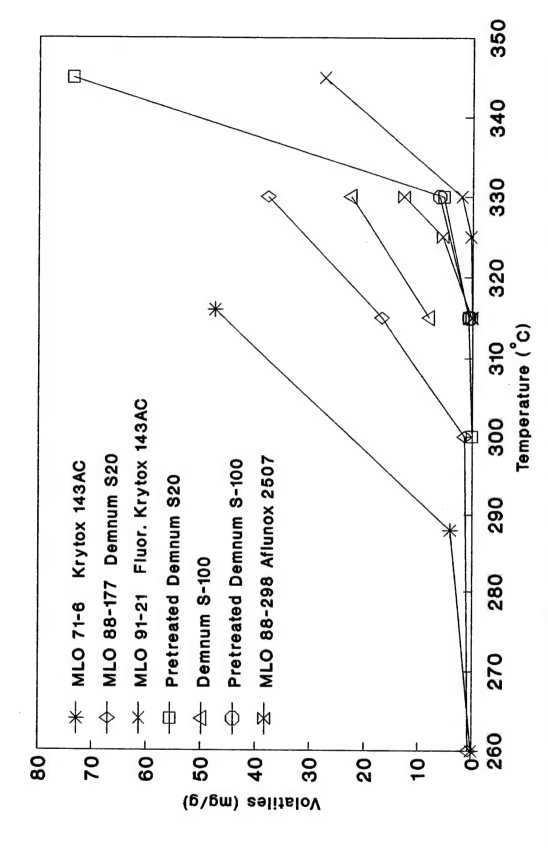
Product of DuPont.

Krytox 143AC which was subjected to fluorination at elevated temperatures.

Product of Daikin Industries.

Demnum S-20 which was exposed to oxygen at 315 and $330\,^{\circ}\mathrm{C}$ in the presence of M-50 alloy for consecutive 24 h periods. (c)

All of these fluids were prepared by direct fluorination of their hydrocarbon precursors. £)



Plots of thermal oxidative degradation data in the presence of M-50 for MLO 71-6, MLO 88-177, MLO 91-21, pretreated MLO 88-177, Demnum S-100, pretreated Demnum S-100 and MLO 88-298. Figure 39.

suspected. In both instances pretreatment at 316-330°C in oxygen, in the presence of M-50 alloy, degrades the low stability chains into volatile products which are then removed. Fluorination provides another way to remove residual hydrogen and was successfully employed in the case of the Krytox 143AC fluid by Exfluor Research Corp.

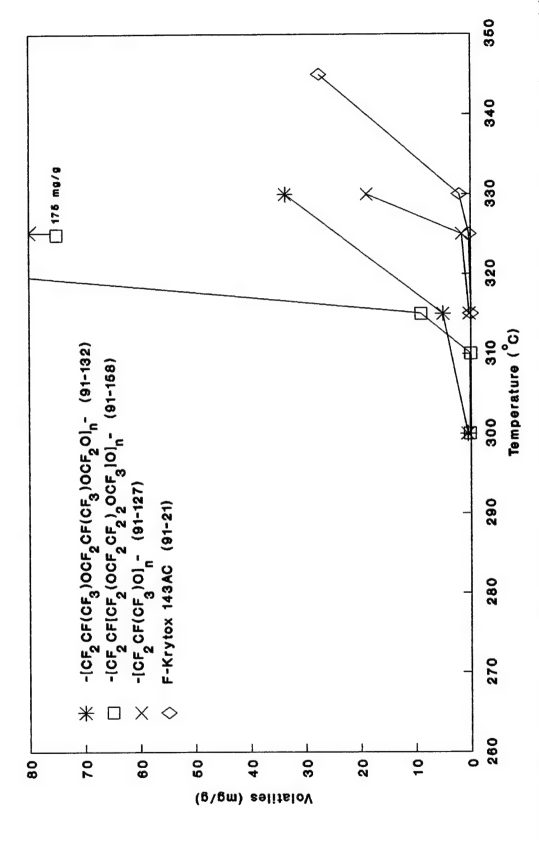
It would appear from the degradation profiles shown in Figure 39 that the inherent thermal oxidative stability of a branched poly(hexafluoropropene oxide) is higher than that of its linear analogue. Specifically, at 315°C the mg/g value for $-[CF(CF_3)CF_2O]_n$ — was found to be 0.03 versus 0.70 for $-(CF_2CF_2CF_2O)_n$ —.

As noted earlier all the other fluids studied were obtained by direct fluorination of their hydrocarbon precursors. No difference was observed in the degradation onset of hydrogen free -[CF(CF₃)CF₂O]_n- prepared by hexafluoropropene oxide telomerization, and of the compound obtained by direct fluorination of poly(propylene oxide) (Figure 40). This confirms that stability is inherent in a given structure and proves the validity of the experimental approach. It is the weakest link or unit which determines the stability of the given material. This is illustrated by the comparison of the degradation onset of 310° C, determined for -(CF₂CF₂O)_n-, and for -[CF₂CF(CF₂OCF₂CF₂CF₂CCF₃OCF₃O]_n-, as shown in Figures 40 and 41. In the second material the sequence of tetrafluoroethylene oxide groups in the branch provides the instability site.

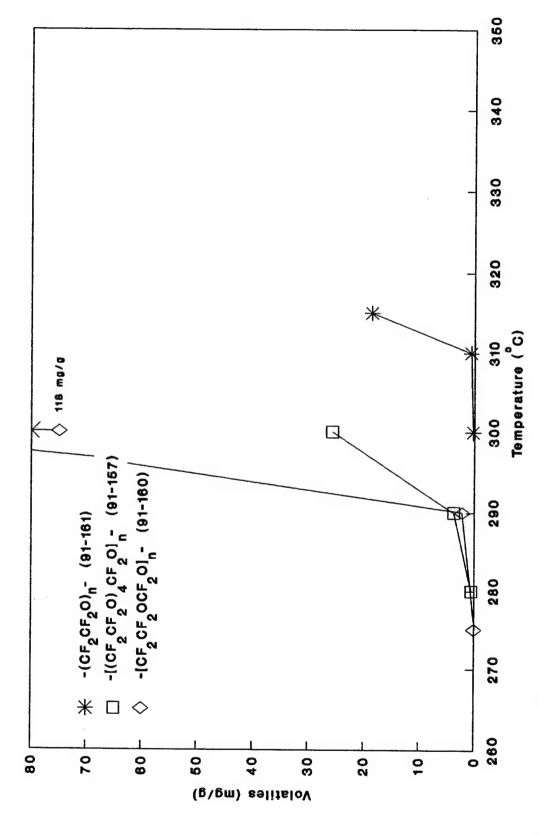
The fluorinated Pluronic L-31 was shown to undergo drastic decomposition between 315 and 330°C (see Figure 42). These results when compared to those obtained for (-CF2CF2O-)n, for which at 315°C the extent of degradation was 18 mg/g (see Figure 40), tend to indicate that adjacent -CF2CF2O- units are absent in this material. The molecular weight of 2800 [see Table 8], the 1:8 ratio of CF2CF2O to CF2CF(CF3)O segments based on ¹⁹F NMR and the method of preparation, (wherein a block of CF(CF3)CF2O segments is terminated by reaction with ethylene oxide), imply structure CF3OCF2CF2O[CF(CF3)CF2O]16CF2CF2OCF3. The material most likely consists of a telomer mixture; however, the general structure must be predominantly CF3OCF2CF2O[CF(CF3)CF2O]xCF2CF2OCF3.

With respect to the fluorinated polyglycol, P15-200, the plot in Figure 42 is in a full agreement with the $^{19}\mathrm{F}$ NMR which gives the ratio of CF(CF₃)CF₂O to CF₂CF₂O segments of 1:1.8 and shows clearly the presence of adjacent CF₂CF₂O units. The latter is further supported by the molecular weight of 4250. Yet, the material degrades less rapidly at 315°C, by a factor of 8, than CF₃O(CF₂CF₂O)_nCF₃, by a factor of 8, based on volatiles formed. Due to its structure, namely the presence of three terminal CF₂CF₂OCF₃ groups per molecule, one would expect this material to undergo degradation at a faster rate than CF₃O(CF₂CF₂O)_nCF₃.

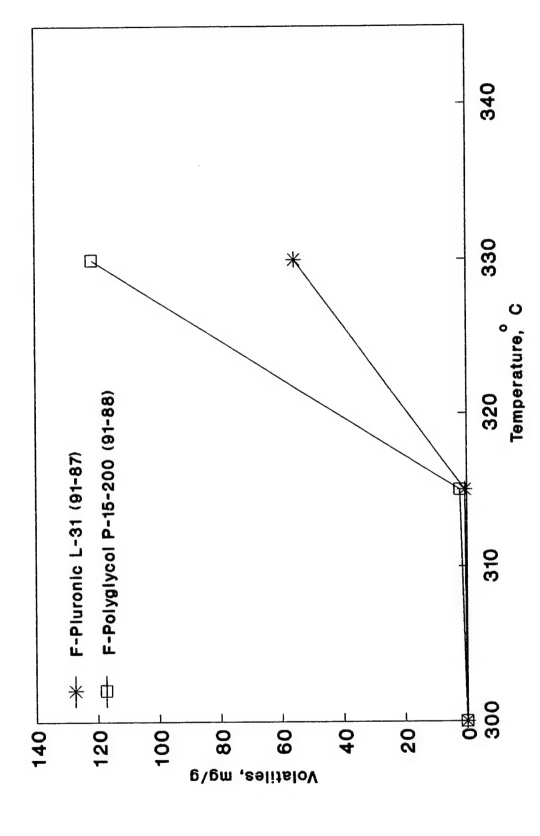
Introduction of a difluoroformyl group into the octafluorobutene oxide system, $-(CF_2CF_2CF_2CF_2O)_n$ -, to give the $-(CF_2CF_2CF_2CF_2CF_2OCF_2O)_n$ - structure, lowers the degradation onset by



Effect of the presence of difluoroformyl groups and tetrafluoroethylene oxide branching on the thermal oxidative stability of poly(hexafluoropropene oxide). Figure 40.



Effect of the presence and relative proportions of difluoroformyl groups on the thermal oxidative stability of poly(tetrafluoroethylene oxide) fluids. Figure 41.

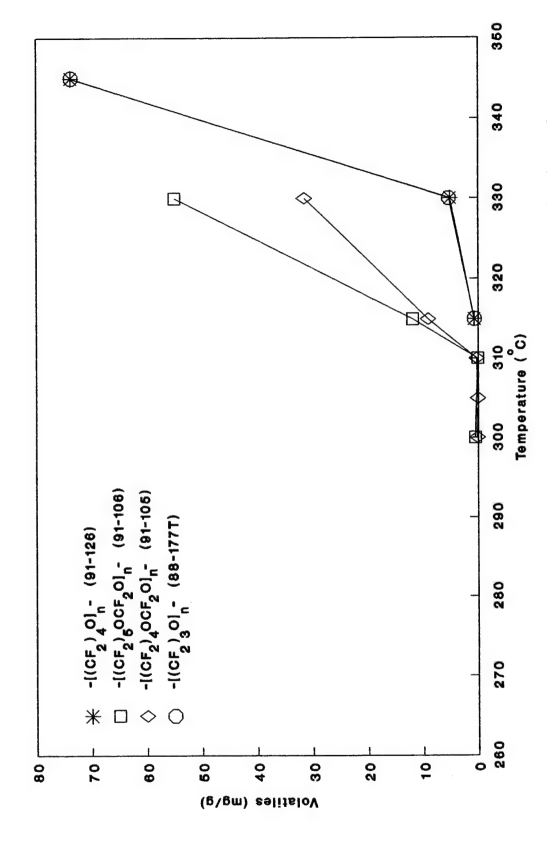


Plots of thermal oxidative degradation data for MLO 91-87 and 91-88 experimental fluids. Figure 42.

5°C, from 315°C to 310°C, as shown in Figure 43. An increase in the number of adjacent CF_2 groups from 4 to 5 has no effect on the degradation onset as is evident from Figure 43. It is of interest that the $-(CF_2CF_2CF_2O)_n$ — and $-(CF_2CF_2CF_2CF_2O)_n$ —arrangements exhibit essentially identical degradation profiles (Figure 43). Apparently, it requires at least three adjacent CF_2 groups to inhibit the "activity" effect of oxygen. Studies performed on aromatic/perfluoroalkylether compounds support these observations [Ref. 22].

It is unexpected that the introduction of difluoroformyl groups into the hexafluoropropene oxide system decreases the degradation onset by 25°C (see Figure 40) versus 5°C for the octafluorobutene oxide structure. Furthermore, the actual onset temperature (300°C) is lower for the copolymer $- [CF_2CF(CF_3) OCF_2CF(CF_3) OCF_2O]_n - \text{ than the } 310°C \text{ determined for } - (CF_2CF_2CF_2CF_2OCF_2O)_n - \text{ (Figure 43)}. It is possible that the presence of the <math>-OCF_2$ - unit disrupts the protective shielding of trifluoromethyl groups and the arrangement approximates more closely a two carbon environment as represented by e.g. poly(tetrafluoroethylene oxide).

The presence of difluoroformyl groups in the tetrafluoroethylene oxide systems results in a drastic drop of the degradation onset from 310°C to 280-285°C. The degradation onsets for the two materials investigated -(CF₂CF₂OCF₂O)_n- and -[(CF₂CF₂O)₄CF₂O]_n- are virtually identical. However, the subsequent rate of degradation above this threshold is



Effect of difluoroformyl groups on thermal oxidative behavior of perfluoropolyalkylethers containing 4 and 5 adjacent difluoromethylene groups. Figure 43.

significantly higher for the composition having the higher diffuoroformyl proportion as is evident from Figure 41.

3.7 DEVELOPMENT OF QUANTITATIVE STRUCTURE/ACTIVITY (THERMAL OXIDATIVE STABILITY IN THE PRESENCE OF M-50) RELATIONSHIPS

Based on earlier data [Ref. 8] and the current investigations, it is obvious that there is a definite interdependence between structure and the thermal oxidative stability of perfluoroalkylethers. Establishment of QSARs to be utilized in conjunction with the quantitative structure/viscosity relationships discussed in Section 3.5 should allow to identify the best perfluoropolyalkylether arrangement for a given application without having to synthesize and evaluate a series of potential candidates. A structure/stability correlation based on energy parameters provides a meaningful approach. Although, as mentioned earlier, the perfluoropolyalkylethers studied are mixtures of telomers, the use of unimolecular compositions in the computational treatments seemed reasonable in view of the results obtained in the structure/viscosity studies. The compositions listed in Table 14 were utilized in this study.

A catalytic process, which is the case for the M-50 promoted degradations, must be dependent on juxtaposition of units or segments in a given material. This is of a particular importance in high molecular weight systems where in a single molecule a number of different configurations can exist. To permit the various molecular orientations, freedom of motion is necessary. In a computational treatment this can be achieved by

imparting energy (velocity) to molecular conformations (obtained using molecular mechanics). Arbitrarily, we have selected 313°K as the equilibrium temperature to determine the various energy components, given in Table 15, which were computed after geometry optimizations. These are average values of four different conformations. Representative conformations of the perfluoropolyalkylethers, listed in Table 15, are presented in Figures 44-48. The lack of uniformity in the structural arrangements illustrates the points discussed above.

To develop QSAR expressions the energy components given in Table 15 were employed in multivariate linear regression analyses. The equations, derived using different energy parameters, together with the corresponding Pearson correlation coefficients (r) are presented in Table 16. Employing just three descriptors gives r of 0.977 and s (standard error of estimate) of 3.61. The graphical presentation is depicted in Figure 49. To determine the validity of this approach, equations using the three energy components, namely bending, Van der Waals and stretch-bend energies, were derived employing 9 out of the 10 fluids. The degradation temperature of the excluded fluid was then calculated. This operation was performed for each of the 10 The relevant data are compiled in Table 17. fluids. agreement between calculated and actual degradation temperature is illustrated in Figure 50.

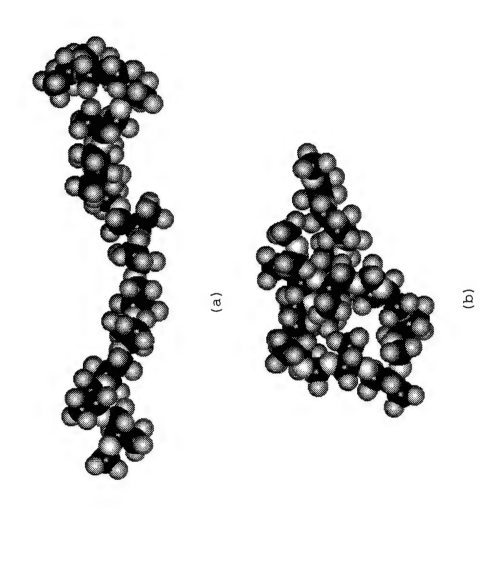
Based on these results and the developed viscosity/structure correlations, apparently it is feasible to

TABLE 15

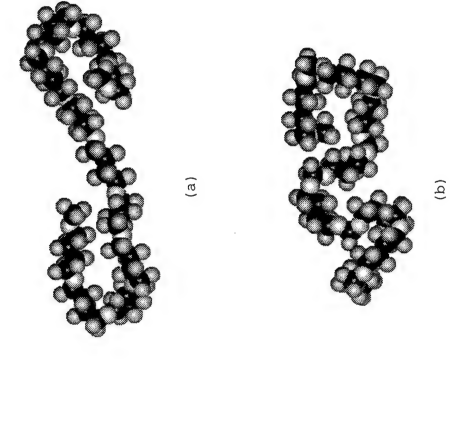
PERFLUOROPOLYALKYLETHER ENERGY COMPONENTS LISTING

		Energ	Energy Components ^a , (kcal/mol)	entsa, (kcal/m	01)
Structure	BOND	BEND	DIHED	VDW	SB	ES
$\mathtt{CF}_3\mathtt{O}$ [($\mathtt{CF}_2\mathtt{CF}_2\mathtt{O}$) $\mathtt{4CF}_2\mathtt{O}$] $\mathtt{4CF}_3$	12.65	13.75	-114.6	-10.17	1,73	226.9
$\mathtt{CF}_3\mathtt{O}\left(\mathtt{CF}_2\mathtt{CF}_2\mathtt{OCF}_2\mathtt{O}\right)$	10.20	14.66	-106.9	-5.87	1.61	189.7
$\mathtt{CF}_3\mathtt{O}$ [$\mathtt{CF}_2\mathtt{CF}$ (\mathtt{CF}_3) $\mathtt{OCF}_2\mathtt{O}$] $_6\mathtt{CF}_3$	18.09	21.41	-104.0	-23.13	2.57	270.1
$CF_3O(CF_2CF_2O)_{20}CF_3$	15.66	15.16	-135.7	0.02	2.04	284.9
CF30[CF2CF(CF2OCF2CF2OCF2CF3)0]5CF3	14.77	23.08	-101.2	-14.01	2.16	244.3
CF3O (CF2CF2CF2OCF2O) 8CF3	21.99	17.03	-125.1	-9.29	2.48	324.9
CF3O (CF2CF2CF2CF2OCF2O) 7CF3	25.93	18.04	-133.4	-12.64	2.90	373.2
$\mathtt{CF}_3\mathtt{O}\left(\mathtt{CF}_2\mathtt{CF}_2\mathtt{CF}_2\mathtt{O}\right)_{14}\mathtt{CF}_3$	24.58	17.04	-145.3	-0.97	2.34	382.0
$\mathtt{CF_3O}(\mathtt{CF_2CF_2CF_2O})_{11}\mathtt{CF_3}$	29.80	18.35	-156.4	-6.68	2.88	442.8
$CF_3O[CF_2CF(CF_3)O]_14CF_3$	20.41	21.12	-116.9	-5.24	2.54	316.8

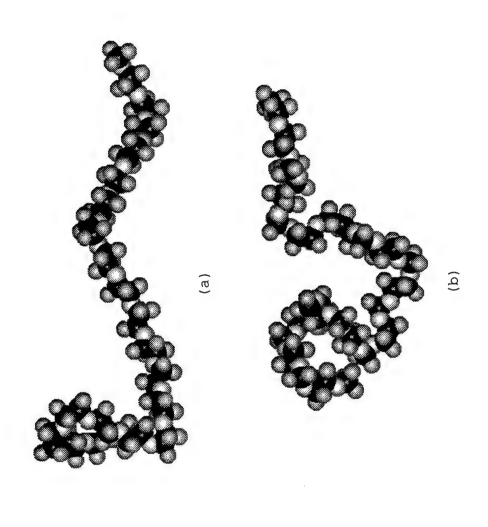
BOND, bonding energy; BEND, bending energy; DIHED, torsion energy; VDW, Van der Waals energy; SB, stretch-bend energy; ES, electrostatic energy. a)



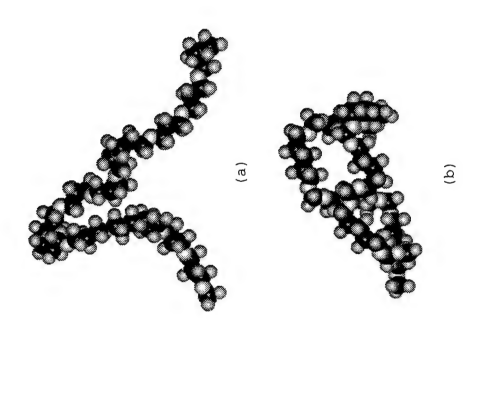
Geometry optimized conformations after molecular dynamics at 40°C: (a) CF30[CF2CF(CF3)0]14CF3, (b) CF30[CF2CF(CF3)0CF2CF(CF3)0CF20]6CF3. Figure 44.



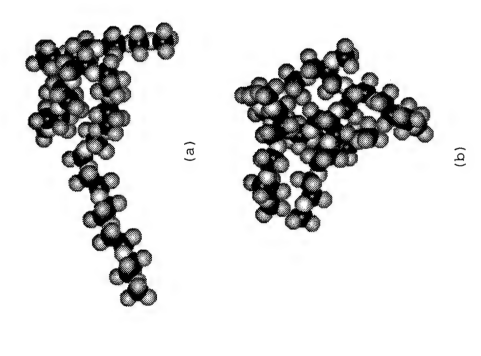
Geometry optimized conformations after molecular dynamics at 40°C: (a) CF30(CF2CF2CF2CF2O) 8CF3. Figure 45.



Geometry optimized conformations after molecular dynamics at 40°C: (a) CF30(CF2CF20)13CF3. Figure 46.



Geometry optimized conformations after molecular dynamics at 40°C: (a) CF30(CF2CF2CF2CF2OCF20)7CF3. Figure 47.



Geometry optimized conformations after molecular dynamics at 40°C: (a) CF30[(CF2CF2OCF2O)4CF3)] (b) CF30[CF2CF(CF2OCF2CF2OCF2OCF3)] 5CF3. Figure 48.

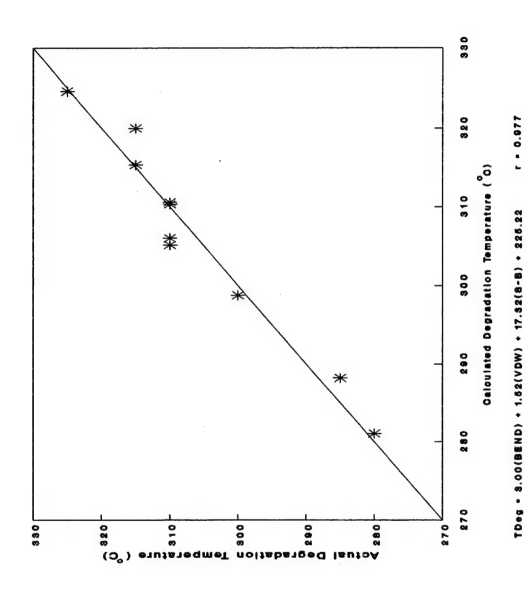
QSAR EXPRESSIONS BASED ON ENERGY DESCRIPTORS TABLE 16

	Constant	231.7	225.4	219.3	225.2	241.8	253.2	242.9
	ES	0.08						
sa	SB	29.98	29.74	30.09	17.32	٠	26.24	18.30
efficient	VDW	1.72	1.75	1.71	1.52	1.56		
Variable Coefficientsa	DIHED	0.16	0.05					
Va	BEND	2.38	2.59	2.68	3.00	4.34		1.18
	BOND	-1.37	-0.70	-0.84	:		÷.	·,
	SC		3.89		3.61	8.05	8.54	10.44
	Rb	0.982	0.982	0.982	0.977	0.860	0.841	0.749
No. of	Variables	9	S	4	8	2	2	2

BOND, bonding energy; BEND, bending energy; DIHED, torsion energy; VDW, Van der Waals energy; SB, stretch-bend energy; ES, Electrostatic energy. a)

R is the Pearson correlation coefficient. S is the standard error.

c)



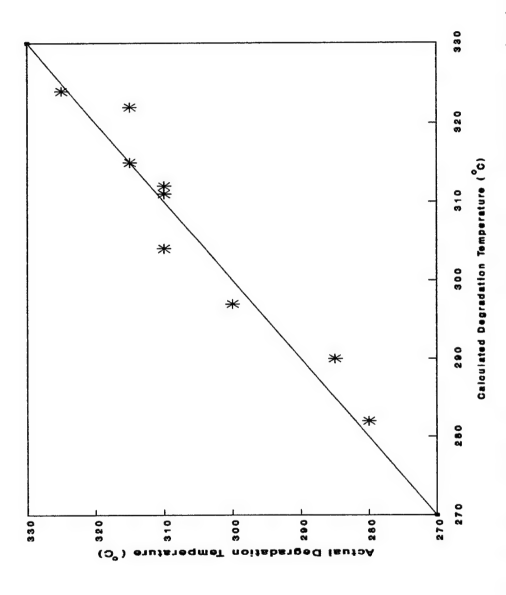
Plot of calculated versus actual degradation temperature as a function of bending, Van der Waals and stretch-bend energies. Figure 49.

TABLE 17

DEGRADATION TEMPERATURES CALCULATED FROM EQUATIONS DERIVED FROM DATA EXCLUDING THE FLUID OF INTEREST

		Va	Variable ^a Coefficients	a ents		Degra Tem	Degradation Temp °C
Fluid excluded	Rb	Bend	VDW	SB	Constant	Calc	Actual
CF30[(CF2CF20)4CF20]4CF3	0.961	2.88	1.47	16.89	228.2	280	282
$\mathtt{CF}_3\mathtt{O}\left(\mathtt{CF}_2\mathtt{CF}_2\mathtt{OCF}_2\mathtt{O}\right)_{13\mathtt{CF}_3}$	0.975	2.93	1.50	15.34	231.5	285	290
CF30[CF2CF(CF3)OCF2CF(CF3)OCF20]6CF3	0.978	3.00	1.62	17.29	225.9	300	297
CF30 (CF2CF20) 20CF3	0.984	3.02	1.41	17.76	222.2	310	304
CF30[CF2CF(CF2OCF2CF2OCF2CF3)0]5CF3	0.977	3.15	1.52	16.62	224.3	310	312
CF30 (CF2CF2CF2OCF20) 8CF3	0.985	3.18	1.55	16.27	224.2	310	304
CF30 (CF2CF2CF2CF20CF20) 7CF3	0.977	2.97	1.51	17.58	225.1	310	311
$\mathtt{CF}_3\mathtt{O}(\mathtt{CF}_2\mathtt{CF}_2\mathtt{CF}_2\mathtt{O})14\mathtt{CF}_3$	0.976	3.00	1.52	17.35	225.2	315	315
$\mathtt{CF}_3\mathtt{O}(\mathtt{CF}_2\mathtt{CF}_2\mathtt{CF}_2\mathtt{O})\mathtt{11}\mathtt{CF}_3$	0.988	2.86	1.57	20.45	221.6	315	322
$CF_3O[CF_2CF(CF_3)O]_14CF_3$	0.970	2.96	1.50	17.34	225.7	325	324

BEND, bending energy; VDW, Van der Waals energy; SB, stretch-bend energy. R is the Pearson correlation coefficient. a) b)



Plot of calculated versus actual degradation temperatures obtained from equations derived excluding the fluid of interest. Figure 50.

design a perfluoropolyalkylether system having specific stability and viscosity characteristics. It must be stressed that, unfortunately, [Ref. 8] perfluoropolyalkylethers exhibiting the highest thermal oxidative stabilities possess also the poorest viscosity/temperature profiles. Thus an optimum composition for a given application will have to be a compromise system.

4. EXPERIMENTAL

General

All solvents used were reagent grade and were dried and distilled prior to use. Operations involving moisture or air sensitive materials were carried out either in an inert atmosphere enclosure (Vacuum Atmospheres Model HE-93BO) under nitrogen bypass or in vacuo. The commercially available starting materials were usually purified by distillation, crystallization, or other appropriate means.

Infrared spectra were recorded as gas spectra, neat (on liquids) and as double mulls (Kel-F oil No. 10 and Nujol) on solids using a Perkin-Elmer Corporation infrared spectrophotometer Model 1330. The mass spectrometric (MS) analyses were obtained employing a Du Pont 21-491B double focusing mass spectrometer attached to a Varian Aerograph Model 2700 gas chromatograph (GC), equipped with a flame ionization detector and a Du Pont 21-094 data acquisition and processing system. The majority of product mixture identifications were performed using combined gas chromatography/mass spectrometry (GC/MS). Gas chromatography was conducted employing a 10 ft x 1/8 in stainless steel column packed with 4% OV-101 on 80/100 mesh Chromosorb G or a 3 ft x 1/8 in stainless steel column packed with 3% Dexsil 300 on 100/120 mesh Chromosorb WAW, using a programming rate of 8°C/minute from 50-300°C. Molecular weights were determined using a Mechrolab Model

302 vapor pressure osmometer. Viscosities were measured with Cannon-Manning semi-micro viscometers.

Materials

Poly(hexafluoropropene oxide), $F(CF(CF_3)CF_2O)_nC_2F_5$, oils of different molecular weights were obtained from Du Pont (Krytox 143 series); the linear isomer, $-[CF_2CF_2CF_2O]_n$ -, was received from Daikin Co. (Demnum S-20 and S-100); Aflunox was procured from PCR Inc. The other perfluoropolyalkylethers were prepared by Exfluor Research Corp. and were provided by the U. S. Air Force.

Degradation

All the tests were carried out in pure oxygen (~400 mm Hg pressure at 25°C) in the presence of an M-50 alloy coupon over 24 h period at the denoted temperatures. At the end of exposure the volatile condensibles were removed in vacuo and weighed. Detailed description of the procedure and apparatus was reported previously [Ref. 23]. The temperature at which the total of volatile condensibles amounted to 0.50 ± 0.25 mg/g (mg of products formed per g of fluid employed) was defined as the degradation onset. To arrive at this point tests were carried out, if necessary, 5°C apart. Thus the onset temperature is given within ± 5 °C.

Computation

The molecular mechanics calculations of energies are based on MM2 developed by Allinger [Ref. 24] modified by Hypercube Inc. (MM+; Hyperchem 3.0 software). The MM+ parameters were

using the U.S. Air Force provided PM3 data set developed by J. J. P. Stewart specifically for perfluoroalkylethers. The bond lengths and angles used are listed below. Bond lengths: C-C = 1.554 Å, C-O = 1.354 Å,

C-F = 1.312 Å. Angles: $C-C-C = 112.2^{\circ}$, $C-C-O = 108.2^{\circ}$, $C-O-C = 124.7^{\circ}$, $O-C-O = 109.4^{\circ}$. To obtain valid derivations, based partially on the average molecular weights of the fluids studied (the exception being Krytox 143AC and its fluorinated product), all the stability computations were carried out on representative telomers having molecular weights as close as feasible to 2500. Each of the structures listed in Table 15 was geometry optimized at 0°K. Using the molecular dynamics program the temperature was increased to 313°K over 3 picoseconds. After equilibrium was reached over 5 picoseconds, data were collected over next 2 picoseconds for a total of 2000 data points. The conformations obtained at 0, 1, 1.5 and 2 picoseconds were geometry optimized and the energy parameters, subsequently calculated at the four time periods, were averaged. These are the energy parameters listed in Table 15. The regression analyses to develop the QSARs were performed with SPSS 4.01 computer program.

Preparation of Cl₂C=CClCCl=CCl(OC₂H₅)

Under nitrogen bypass to stirred hexachlorobutadiene (400 g, 1.53 mol), at $78\,^{\circ}\text{C}$ was added (via an addition funnel) 576 mL of NaOC₂H₅ solution in ethanol (21% by weight) over a period of 6

hours. After the addition was completed, the solution was cooled to room temperature, followed by addition of water (800 mL) and ether extraction. Following separation the organic layers were dried over MgSO₄. Removal of the solvent gave a dark yellow solution (355 g). Based on the GC/MS, it contained 45% of the product which consisted of at least three isomers. Purification using a 3 ft spinning band column gave 95 g (23% yield) of 96% pure product, bp 60-61°C at 0.40-0.42 mm Hg. The two impurities present were the starting material, hexachlorobutadiene, and the di-substituted by-product. A representative MS of Cl₂C=CClCCl=CCl(OC₂H₅) is given in Table 18.

Preparation of Cl3CCC1=CC1COC1

At room temperature, into $Cl_2C=CClCCl=CCl(OC_2H_5)$ (25.9 g, 95.8 mmol) was bubbled chlorine over 7 3/4 hours; subsequently, the mixture was heated under nitrogen bypass at 125°C for 2 hours. The crude product (15.6 g) contained 66% of the desired product, as determined by GC. Purification by distillation, through a short column in vacuo, gave 18.9 g (71.5% yield) of 86% pure $Cl_3CCCl=CClCoCl$, bp 42-43°C at 0.001 mm Hg. The two impurities were $Cl_2C=CClCoCl$ and $Cl_2C=CClCcl=CCl(OC_2H_5)$. The mass spectrum of $Cl_3CCCl=CClCoCl$ is given in Table 19.

Preparation of Hexachlorofuran

To Cl₃CCCl=CClCoCl (58.3 g, 165 mmol) was added FeCl₃ (0.6 g), and the mixture was stirred under nitrogen for 3 hours at 170° C. The product was then distilled out, bp $44-48^{\circ}$ C at 0.001 mm

TABLE 18 $\label{eq:table_problem} \mbox{ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF $$\operatorname{Cl}_2\mathrm{C=CCl-CCl=CCloc}_{2}\mathrm{H}_5$$}$

m/e	%	m/e	%	m/e	8	m/e	8
35	17.8	117	25.0	170	17.4	214	6.0
36	51.5	118	17.5	171	27.4	215	23.9
37	14.7	119	24.9	172	15.0	216	5.0
38	21.5	120	13.9	173	13.9	217	15.8
43	9.9	121	12.8	174	9.0	219	6.4
47	19.6	123	90.9	176	76.2	223	7.0
49	12.1	124	8.8	177	59.2	225	9.4
59	5.2	125	74.3	178	85.3	227	6.6
63	25.3	126	7.0	179	71.1	233	16.0
64	6.3	127	22.8	180	58.7	234	5.8
65	12.9	133	9.7	181	40.7	235	21.6
70	6.2	134	19.5	182	24.6	237	11.9
71	34.4	135	11.6	183	18.0	239	5.0
72	16.0	136	14.8	184	5.0	240	42.8
73	23.9	137	5.5	188	14.9	241	9.5
74	7.0	141	93.3	189	5.1	242	69.6
82	9.8	142	14.4	190	17.7	243	9.8
83	16.6	143	87.8	191	6.0	244	47.1
84	11.3	144	12.6	192	10.5	245	7.7
85	11.2	145	39.0	197	6.5	246	22.9
87	14.3	146	6.0	199	6.0	248	7.2
89	8.6	147	11.6	204	50.4	268	88.4
94	11.2	153	12.9	205	84.1	269	17.1
96	7.8	154	7.7	206	69.9	270	100.0
99	12.0	155	14.2	207	84.2	271	24.0
106	59.0	156	5.2	208	42.5	272	82.4
107	22.9	157	26.7	209	63.1	273	15.5
108	35.4	159	25.6	210	17.4	274	37.7
109	17.9	161	13.6	211	30.9	275	7.1
110	12.3	163	5.8	212	7.2	276	11.6
111	5.8	169	29.1	213	25.9		

Peaks having intensities lower than 5% of the base peak and lower than m/e 31 are not reported.

m/e	ion	m/e	ion
268	М	204	$M - HCl - C_2H_4$
240	$M - C_2H_4$	169	$M - HCl - C_2H_4 - Cl$
233	M - Cl	106	$M - HCl - C_2H_4 - Cl - COCl$

TABLE 19

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF Cl₃C-CCl=CClCOCl

m/e	8	m/e	%	m/e	%	m/e	%
35	31.6	89	18.3	131	6.8	212	18.4
36	54.7	90	8.2	134	5.0	213	100.0
37	19.0	94	18.9	141	62.6	214	29.3
38	17.0	96	13.9	142	7.4	215	88.6
47	25.2	103	5.9	143	59.8	216	26.3
49	18.4	105	11.3	144	6.4	217	67.7
53	14.3	106	78.2	145	24.1	218	11.9
54	9.8	107	17.7	147	8.1	219	19.8
59	11.7	108	52.0	176	21.7	239	44.8
61	5.2	109	6.2	178	21.9	240	5.5
63	50.0	110	18.3	179	8.5	241	61.1
65	22.6	112	6.6	180	17.3	242	8.5
70	19.9	117	68.4	181	6.3	243	44.9
71	59.6	118	6.6	182	7.0	244	5.1
72	19.3	119	61.2	186	9.8	245	20.6
73	22.7	120	6.1	188	9.9	247	5.8
74	5.7	121	25.0	204	15.3	274	19.0
82	18.3	122	5.0	206	15.4	276	24.0
83	5.1	123	13.3	208	11.6	278	21.5
84	15.9	125	6.6	210	5.4	280	13.6
87	13.2	129	6.2	211	95.6	282	5.1
88	13.0						

Peaks having intensities lower than 5% of the base peak and lower than m/e 31 are not reported.

m/e	ion	m/e ion
274	M	$\overline{176}$ \overline{M} – COCl – Cl
239	M - Cl	141 M - COC1 - 2C1
211	M - COCl	117 Cl ₃ C
204	M - 2C1	5230

Hg, giving 43.3 g (94.8% yield) of perchloro-2,5-dihydrofuran, which solidified after cooling below 0°C. Mass spectrum is given in Table 20.

Preparation of 2,2,5,5,-Tetrafluoro-3,4-dichlorofuran from hexachlorofuran

To hexachlorofuran (42.8 g, 0.155 mol) were added SbF₃ (56 g, 0.31 mol) and SbCl₅ (24 g, 80 mmol). The mixture was then stirred at 100°C for 2 hours under nitrogen atmosphere.

Distillation, bp 72-74°C, gave 17.5 g (52% yield) of the desired product (purity >97%); the mass spectrum is presented in Table 21.

Attempted Preparation of 2,2,5,5,-Tetrafluoro-3,4-dichlorofuran from Dichloromaleic Anhydride

In a typical reaction, to a 600 mL stainless steel bomb (Parr Instrument Co.) was added dichloromaleic anhydride (50.0 g, 0.30 mol) followed by anhydrous HF (4 mL). Next SF4 (259 g, 2.4 mol) was condensed in at -78°C using a high pressure hose. Subsequently, the reactor was warmed to room temperature, placed in a sand bath on a shaker, and heated at 345°C for 160 hours with constant shaking. The bomb was then cooled to 0°C (in an ice bath) and vented slowly. This was followed by warming to room temperature, and after 1 hour the contents were poured into a plastic bottle containing 4 g NaF to remove any remaining HF. After transfer to a round bottom flask, the products were separated from NaF/NaHF2 mixture by bulb-to-bulb distillation in vacuo, initially at room temperature, then at 40°C. A 50.5 g

TABLE 20

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF

m/e	%	m/e	%	m/e	8	m/e	%
35	50.9	98	8.5	157	13.1	218	8.1
36	95.7	102	13.7	159	5.0	219	21.6
37	28.8	103	16.5	166	5.9	223	38.2
38	29.1	104	10.5	176	38.8	224	7.2
47	56.3	106	98.5	177	7.3	225	54.7
48	12.1	107	36.6	178	49.1	226	9.4
49	25.8	108	90.2	179	9.0	227	42.0
53	19.5	109	19.6	180	29.4	228	6.9
54	16.1	110	27.9	181	5.4	229	20.0
59	20.2	117	57.3	182	13.1	231	7.2
61	10.1	118	37.4	188	33.3	239	88.0
63	48.8	119	61.5	189	8.1	240	22.2
65	23.5	120	29.9	190	38.2	241	96.7
70	15.2	121	30.4	191	10.8	242	21.9
71	79.0	122	12.5	192	24.5	243	89.8
72	25.3	123	8.7	193	6.2	244	21.0
73	47.5	129	9.0	194	10.9	245	56.7
74	7.9	130	5.3	204	31.9	246	9.1
82	25.5	131	9.0	205	6.2	247	20.1
83	23.8	133	5.0	206	41.4	258	21.3
84	22.2	134	8.2	207	8.9	260	29.8
85	13.6	136	5.3	208	24.5	261	5.4
86	8.0	141	85.7	209	5.9	262	26.4
87	20.1	142	17.7	210	12.5	263	5.0
88	18.1	143	87.0	211	87.3	264	18.2
89	24.8	144	17.8	212	19.8	266	7.7
90	18.3	145	59.4	213	100.0	274	6.8
91	6.9	146	8.1	214	22.0	276	10.8
94	8.5	147	16.5	215	91.3	278	8.7
95	31.7	153	22.5	216	17.1	280	5.4
96	24.1	155	21.9	217	59.3		

Peaks having intensities lower than 5% of the base peak and lower than m/e 31 are not reported.

m/e	ion	m/e	ion	m/e	ion
274		204	\overline{M} - 2Cl	63	COC1
	M - Cl	176	M - COCl ₂	47	CCl
211	M - COCl		$M - COCl_2^2 - Cl$		

TABLE 21

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF

$$\begin{array}{ccc} \text{Clc} & \longrightarrow \text{Ccl} \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ | & | \\ |$$

m/e	ૠ	m/e	જ	m/e	%	m/e	%
31	37.7	71	4.6	94	10.4	163	54.3
32	11.3	72	5.4	96	6.8	165	37.4
35	7.1	73	4.6	97	5.3	167	6.7
36	7.8	74	41.2	109	100.0	175	25.3
47	26.9	78	6.3	110	7.7	177	9.0
49	5.6	85	10.2	111	47.3	182	4.1
50	6.1	87	7.1	144	11.1	191	5.5
55	21.4	90	14.2	146	7.3	193	3.3
59	8.9	92	4.4	147	12.5	210	21.1
61	3.3	93	12.1	149	3.7	212	14.3
69	46.6						

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

Significant Ions in Support of Structure and Composition

m/e	<u>ion</u>
210	М
175	M - Cl
163	M - CCl
147	M - COCl
109	M - COF ₂ -Cl

mixture consisting of 15% of the desired furan, and 84% lactone was collected.

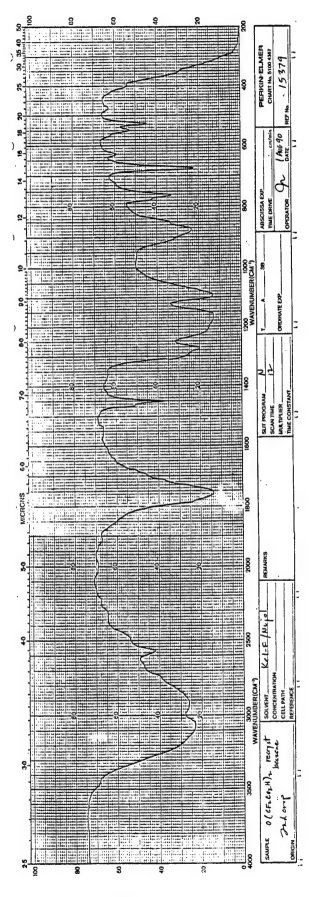
Treatment of the Lactone/Furan Mixture with SF4

The 50.5 g mixture obtained from the reaction of dichloromaleic anhydride with SF₄ was retreated with SF₄ (98 g, 0.91 mol) at 345°C over 160 hours using the procedure described above. After workup 47 g of the crude product was obtained which consisted of 74% 2,2,5,5-tetrafluoro-3,4-dichlorofuran, 23% lactone and 1.5% CF₃CCl=CClCF₃ and CF₃CCl=CClCoF. The furan, bp 72-74°C (purity 97%, GC), was isolated by distillation.

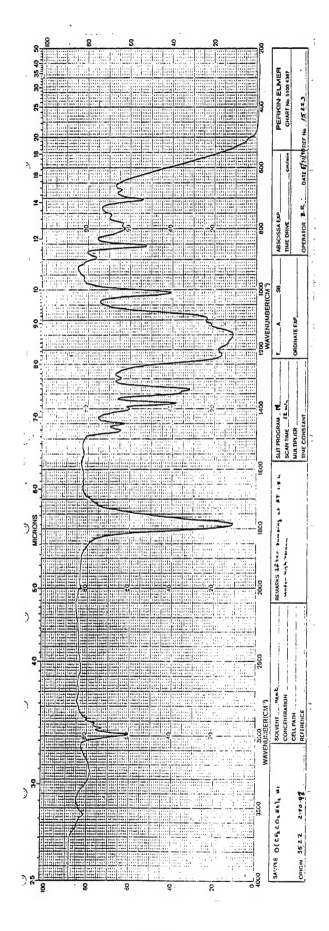
Preparation of Perfluorooxydiacetic acid

In a typical reaction, a 250 mL round bottom flask equipped with a magnetic stir bar, a Claisen adapter attached to a condenser and additional funnel, was charged with KOH (10.7 g, 190 mmol), KMnO4 (15.0 g, 94.8 mmol), and 85 mL distilled water.

After stirring for 10 minutes (until all KOH dissolved) 2,2,5,5-tetrafluoro-3,4-dichlorofuran (10.0 g, 47.4 mmol) was added dropwise over a period of 15 minutes. The reaction mixture was then stirred for 1 hour at room temperature. Subsequently, the temperature was slowly raised to 44°C and held there for 18 hours. After cooling to room temperature, H₂SO₄ (50 mL, 75%) was added slowly over a 30 minute period. The entire reaction mixture was continuously extracted with diethyl ether over a 7 hour period. The ether extracts were dried over MgSO₄; ether was removed by distillation under nitrogen bypass to give 7.5 g of crude product.



Infrared spectrum (Kel-F/Nujol double mull) of perfluoro-oxydiacetic acid (IR# 15379 ID# 2-90-140). Figure 51.



Infrared spectrum of $\rm C_2H_5O_2CCF_2OCF_2CO_2C_2H_5$ (ID# 2-90-97, IR# 15223), thin film. Figure 52.

m/e	%	m/e	%	m/e	%	m/e	%
27	52.7	55	6.8	94	9.5	162	10.9
28	68.3	56	5.7	95	30.2	187	14.9
29	100.0	57	7.2	96	53.9	189	26.7
30	44.9	59	40.2	97	12.1	190	17.8
31	45.2	60	8.5	98	5.1	191	5.6
32	8.1	65	10.5	115	11.8	207	21.6
33	6.4	66	10.1	116	9.5	210	5.6
41	11.8	67	10.2	121	10.1	215	30.7
42	11.0	73	22.1	123	35.8	216	7.1
43	34.0	74	17.8	124	49.8	217	9.3
44	34.8	75	7.0	125	31.5	225	21.5
45	51.3	76	6.4	126	6.1	235	31.5
46	9.5	77	10.1	139	6.0	236	7.6
47	33.2	78	28.6	142	10.5	243	17.1
48	5.1	79	11.8	152	23.8	263	44.6
50	30.7	80	11.4	153	5.0	264	20.4
51	38.3	93	14.3	161	10.7	291	5.0

Peaks having intensities less than 5% of the base peak and lower than m/e 27 are not reported.

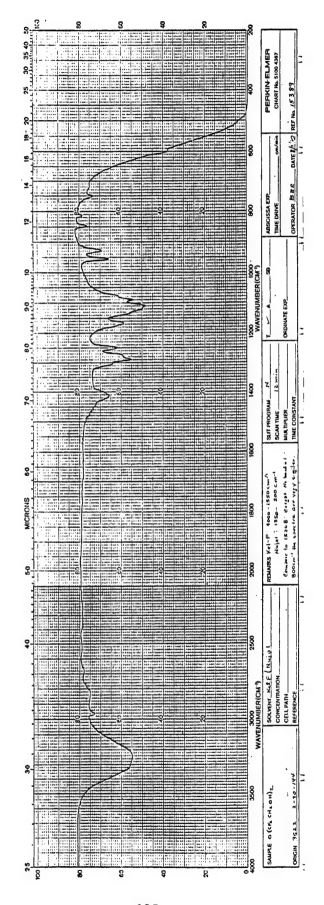
Recrystallization from hot benzene in an inert atmosphere enclosure (the material is extremely hygroscopic) gave pure perfluorooxydiacetic acid, 5.9 g (61% yield) mp 77-78°C. Its infrared spectrum is given in Figure 51.

Preparation of Diethyl Ester of Perfluorooxydiacetic Acid

A 100 mL round bottom flask equipped with a magnetic stir bar was charged with ethanol (10 mL), benzene (50 mL), concentrated H₂SO₄ (1 mL), and perfluorooxydiacetic acid (2.3 g, 11.2 mmol). The flask was fitted with a reflux condenser and the mixture was refluxed overnight under nitrogen bypass. After cooling to room temperature it was poured into 50 mL of distilled water. The bottom layer was removed and extracted with 3 x 25 mL portions of diethyl ether. The organic portions were combined, washed with water, and dried over MgSO₄, and the volatile solvents were removed by distillation to give the ethyl ester, 2.2 g (75% yield; 83% purity, GC). The infrared spectrum is given in Figure 52, the mass spectrum in Table 22.

Preparation of HOCH2CF2OCF2CH2OH

In the inert atmosphere enclosure, a 100 mL round bottom three necked flask was charged with LiAlH4 (1.2 g, 32 mmol) and fitted with a reflux condenser, nitrogen bypass valve, and a 25 mL addition funnel. Under nitrogen bypass, diethyl ether was added slowly via the addition funnel. The mixture was cooled to 0°C (ice/water bath) and a solution of EtO2CCF2CCF2CO2Et (2.0 g, 7.63 mmol) in diethyl ether (20 mL) was added dropwise over a 40 minute



of HOCH2CF2OCF2CH2OH (IR# 15389, Infrared spectrum (Kel-F/Nujol double mull) ID# 2-90-144). Figure 53.

period. The reaction was instantaneous, giving off gases after each drop. The mixture was allowed to warm slowly to room temperature, followed by 1 hour reflux. The excess of LiAlH₄ was destroyed with ethyl acetate at 0°C. After 15 minutes of stirring, water (5 mL) was added followed by aqueous H₂SO₄ (100 mL, 20%). The bottom layer was removed and continuously extracted with diethyl ether for 6 hours. The original ether layer was combined with the ether extract, dried over MgSO₄, and following solvent removal 1.3 g (96% yield) of crude diol was obtained. Crystallization from benzene gave white needles of HOCH₂CF₂OCF₂CH₂OH, 1.1 g (80.8% yield) mp 68-69°C (literature 68°C, Ref. 13); the infrared spectrum is given in Figure 53; the mass spectra of the free alcohol and of its trimethylsilyl derivative are given in Tables 23 and 24 respectively.

Attempted Preparation of O(CF2COC1)2

In an inert atmosphere enclosure, a 100 mL flask containing a magnetic stir bar was charged with 2.5 g (12.1 mmol) of $O(CF_2CO_2H)_2$ and 6.2 g (30.3 mmol) of phthalolyl chloride. The flask was then fitted with a reflux condenser and heated at 160°C for 2 hours under nitrogen bypass. After cooling to room temperature the volatiles were removed in vacuo initially at room temperature then at 50-55°C and were collected in a -196°C cooled trap.

The volatiles (2.4 g, 81% yield) exhibited 39 mm Hg vapor pressure at 24.1°C. Analysis by GC and GC/MS found the material

TABLE 23

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF HOCH2CF2OCF2CH2OH

m/e	8	m/e	8	m/e	8	m/e	8
14	15.4	48	5.4	77	7.5	119	5.6
15	6.6	49	17.2	78	24.2	122	5.0
18	9.6	50	15.4	79	18.6	123	7.0
19	12.6	51	67.6	80	64.1	125	3.2
20	13.4	52	7.7	81	81.4	127	63.9
26	5.4	53	9.6	82	31.5	128	28.1
27	4.7	54	5.0	83	11.8	129	5.1
28	38.0	58	15.1	91	13.9	130	5.3
29	65.6	59	19.7	92	47.7	131	3.2
30	39.7	60	18.3	93	14.6	138	7.6
31	100.0	61	74.1	94	9.3	139	55.1
32	45.2	62	53.9	95	3.0	140	10.5
33	97.1	63	30.3	97	7.0	141	18.5
34	9.9	64	69.0	98	3.9	142	3.5
36	3.8	65	46.6	99	4.1	143	7.1
41	5.7	66	19.0	101	13.8	145	3.5
42	59.0	67	40.1	108	14.9	147	15.2
43	25.1	69	3.0	109	6.8	148	8.0
44	17.2	71	14.0	110	10.0	158	5.7
45	27.6	72	5.1	111	36.4	161	3.9
46	16.2	73	4.0	112	7.4	178	1.1
47	34.1	74	5.5	113	16.0	179	6.7

Peaks having intensities lower than 3% of the base peak and lower than m/e 14 are not reported.

<u>m/e</u>	<u>Intensity</u>	<u>Assignment</u>	m/e	<u>Intensity</u>	<u>Assignment</u>
178	1.1	M^+	94	9.3	(COF) 2 ⁺
161	3.9	$(M-OH)^+$	81	81.4	$(CF_2CH_2OH)^+$
147	15.2	(M-CH2OH) +	64	69.0	$(CF_2CH_2)^+$
141	18.5	$(M-OH-HF)^+$	61	74.1	(CF=CHOH) +
105	50.0	(31	100.0	(CH ₂ OH) +
127	63.9	(CFOCH ₂ CF ₂) ⁺	20	13.4	(HF) ⁺

TABLE 24

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF

Me₃SiOCH₂CF₂OCF₂CH₂OSiMe₃

m/e	%	m/e	%	m/e	%	m/e	%
15	12.3	51	7.1	80	6.1	139	4.9
27	9.2	55	7.2	81	25.7	151	11.6
28	18.7	57	9.9	82	3.8	152	5.6
29	36.5	58	15.3	83	3.6	153	35.8
30	6.2	59	19.0	88	7.1	154	11.1
31	16.7	60	3.6	91	27.2	155	6.7
32	3.0	61	33.9	92	5.1	157	17.8
33	20.1	62	7.2	93	4.7	158	3.4
34	3.1	63	33.9	103	32.4	169	4.3
35	6.2	64	13.5	104	10.2	187	14.8
42	17.3	71	4.2	105	4.9	211	7.3
43	28.8	72	15.2	107	54.4	215	27.8
44	17.9	73	100.0	108	11.9	216	8.4
45	37.3	74	34.9	109	7.9	217	4.1
46	5.4	75	28.9	111	8.4	227	14.4
47	26.8	76	11.6	119	6.1	228	4.6
48	3.3	77	85.1	135	50.8	307	9.5
49	23.6	78	19.6	136	11.8		
50	3.0	79	16.1	137	8.5		

Peaks having intensities lower than 3% of the base peak and lower than m/e 15 are not reported.

m/e	<u>m/e</u>
$307 - [M - CH_3]^+$	103 - $Me_3SioCH_2^+$
153 - Me ₃ SiOCH ₂ CF ₂ +	$73 - Me_3Si^+$

to consist of 16% perfluorooxydiacetic anhydride, 61% diacid chloride, and some unidentified higher boiling fractions.

Preparation of Methyl Ester of Perfluorooxydiacetic Acid

bar was charged with 3.3 g of a mixture containing 79% of perfluorooxydiacetic anhydride and 7.1% diacyl chloride (based on GC area %). After cooling to 0°C, 15 mL of methanol were added dropwise via addition funnel. A vigorous reaction occurred with the evolution of white fumes. The addition was completed in 0.5 hour. At this stage, 0.5 mL of concentrated H₂SO₄ was added and the mixture heated at 78°C for 19 hours. Following cooling to room temperature the mixture was treated with diethyl ether (75 mL) and then washed with 3 x 25 mL portions of water. The organic layer was dried over anhydrous MgSO₄ to give after solvent removal 3.07 g (94.2% yield) of the desired methyl diester. Its mass spectrum is presented in Table 25 and the infrared spectrum in Figure 54.

Preparation of Oxalyl Fluoride

In an inert atmosphere enclosure a 300 mL three-neck round bottom flask equipped with a magnetic stirring bar, reflux condenser, addition funnel, and nitrogen bypass inlet was charged with sodium fluoride (45 g, 1.07 mol, dried at 250°C in vacuo) followed by acetonitrile (90 mL, distilled from P2O5). Oxalyl chloride (45.4 g, 0.358 mol) was weighed into the addition funnel. After attachment to nitrogen bypass and connection to a receiver

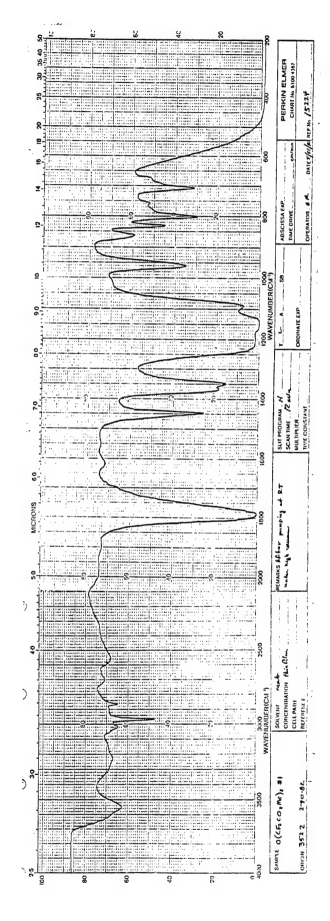


Figure 54. Infrared spectrum of ${
m CH_3O_2CCF_2OCF_2CO_2CH_3}$ (ID# 2-90-82, IR# 15224), thin film.

TABLE 25

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF CH3OC(0)CF2OCF2C(0)OCH3

m/e	%	m/e	%	m/e	%	m/e	%
31	35.1	62	5.9	90	9.2	126	4.7
32	12.4	64	5.2	93	15.2	143	8.3
33	8.0	65	61.5	94	33.0	156	8.5
43	26.8	66	20.0	95	8.0	175	41.7
44	23.5	67	8.0	96	2.6	176	7.2
45	80.3	69	5.3	108	4.2	207	5.3
46	11.3	73	4.2	109	72.2	215	8.6
47	36.5	74	4.1	110	19.7	235	3.3
49	3.0	75	14.4	111	4.4	249	25.8
50	39.2	78	26.8	113	3.0	250	5.7
51	24.8	79	20.8	116	7.1	343	3.1
59	100.0	80	13.2	121	17.0		
60	22.5	81	61.2	124	83.9		
61	9.7	82	8.4	125	16.4		

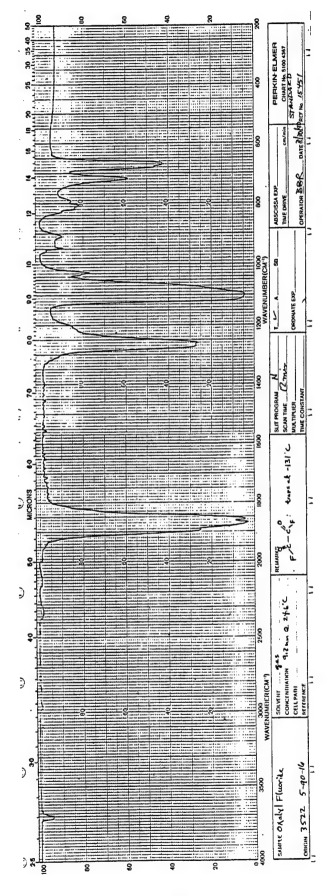
Peaks having intensities less than 3% of the base peak and lower than m/e 31 are not reported.

at -78°C, oxalyl chloride was added to the NaF/CH₃CN mixture over a period of 30 minutes at room temperature, followed by stirring for an additional hour. Subsequently, the reaction flask was heated at 75-85°C for 13 hours. After heating, the receiver flask was closed off and attached to the vacuum system and the products fractionated through traps held at -63°, -111°, and -196°C. Several fractionations were necessary to collect pure oxalyl fluoride in the -111°C trap (17.54 g, 52% yield). The infrared spectrum is given in Figure 55.

Direct Fluorination of Tetraglyme and Preliminary Analyses by Allied-Signal

An 8 hour reaction of tetraglyme was run in a continuous flow aerosol reactor producing 10.7 g of crude product. The reaction conditions were as follows: reactor temperature -20°C, fluorine:hydrogen ratio 1.4:1, tetraglyme pump flow rate 0.5 cc/minute, photochemical stage was on. Under the conditions of the test, only a small portion of the sample is actually introduced into the reactor through a small orifice. The rest is returned to the reservoir. The material is injected into the reactor as a fine spray. Any aerosol reactor is characterized by a long reaction time and a very low through-put.

After completion of the reaction, the product was fractionated on a vacuum line using a three-stage process of -45° C, -78° C, and -196° C. The bulk of the sample stayed in the -45° C trap, indicating very little fragmentation of the tetraglyme



Infrared spectrum of oxalyl fluoride, FC(0)C(0)F (IR# 15451, ID# 5-90-16; gas phase: 9.2 mm at 24.6°C). Figure 55.

during the reaction. The contents of the -45° C trap were then fractionated again through two traps at -5° C and -45° C to remove any underfluorinated tetraglyme. The -45° C fraction which amounted $40-45^{\circ}$ of the product, showed no residual hydrogen left using infrared analysis. The -5° C fraction exhibited a weak absorbance at 3000 cm⁻¹, indicating residual C-H bonds. GC analysis of the -45° C trap showed the sample to contain three peaks:

Retention Time	Peak Area	Area %
2.60 minutes	884	2.269
2.96 minutes	653	1.676
4.11 minutes	37423	96.05

The column used was a 12 ft x 1/8 in 5% SP-2401 on 100/120 Supelcoport. The temperature profile was 34° C for 2 minutes and ramped at 10° C/minute until all peaks eluted.

19F NMR of the sample gave three peaks at -56.548 ppm (t), 89.813 ppm (s), and 91.802 ppm (q) referenced to CFCl₃. The NMR spectrum correlates very well with literature values. Very little contamination of the sample was seen in the spectrum, indicating that most of the impurities in the GC were from other perfluorinated tetraglyme fragments. Proton NMR was run to determine hydrogen content; however, it showed no evidence of residual hydrogen in the sample.

GC/MS and Infrared Spectral Analyses Performed by Ultrasystems on Perfluorotetraglyme Samples Received from Allied-Signal and NASA Lewis

In Figure 57 are given the gas chromatograms run on two different columns (OV-101 and Porapak Q) by Ultrasystems on the sample of perfluorotetraglyme received from Allied-Signal; in Figure 56 are given the corresponding GC traces of a perfluorotetraglyme sample obtained from NASA Lewis on 3 July 1985. For the samples run on the OV-101 column (Figures 56a and 57a), the purity of the material obtained from NASA (prepared by Exfluor Research Corp.) is definitely higher, 95 vs. 89%.

For GC/MS analysis the two samples were run on the Porapak Q column (see Figures 56b and 57b) in order to better separate the major impurity peaks which elute before the perfluorotetraglyme. For NASA Lewis material, the major impurity (6.3 minutes reaction time (RT)) appears to be CF3CF2O(CF2CF2O)3CF3; the Allied-Signal sample also contains this impurity (6.0 minutes RT) in addition to CF3O(CF2CF2O)3CF3 (4.3 minutes RT). The mass spectrum of the 6 minute peak in the Allied-Signal material also exhibited a high output at m/e 59 which does not appear in the NASA Lewis sample. In fact, none of the perfluorinated ethers, including the perfluorotetraglyme (MS, Table 26), show any significant output at m/e 59. The m/e 59 ion is generally associated with [CO2CH3]+; [C2FO]+ and [C3H4F]+ can also account for this ion. The presence of m/e 15 [CH3]+ in the mass spectrum, however, seems to support [CO2CH3]+ for the m/e 59

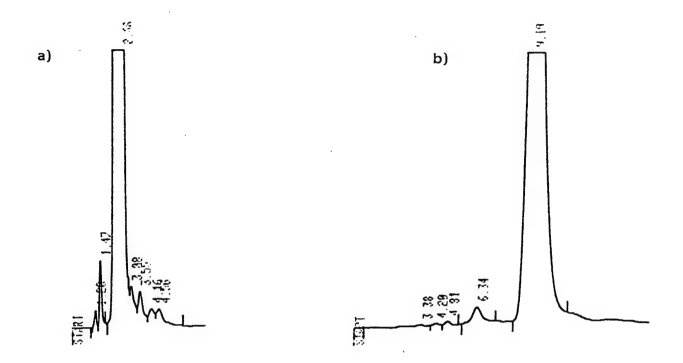


Figure 56. Gas chromatograms of $CF_3O(CF_2CF_2O)_4CF_3$ received from NASA Lewis: (a) 10' x 1/8" SS column, 4% OV-101 on Chromosorb G, isothermal 35°C; (b) 8' x 1/8" SS column, Porapak Q, programmed from 200-220°C at 8°/min.

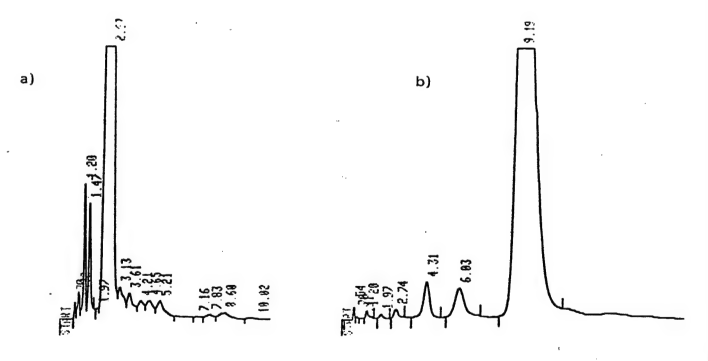


Figure 57. Gas chromatograms of $CF_3O(CF_2CF_2O)_4CF_3$ prepared by Allied-Signal: (a) $10^1 \times 1/8^{11}$ SS column, 4% OV-101 on Chromosorb G, isothermal 35°C; (b) $8^1 \times 1/8^{11}$ SS column, Porapak Q, programmed from 200-220°C at 8° /min.

TABLE 26

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF

CF₃O(CF₂CF₂O)₄CF₃ (MW 618)

m/e	8	m/e	8	m/e	8	m/e	8
28	14.2	78	16.2	136	10.6	252	3.3
31	22.1	81	6.9	163	30.7	279	12.5
47	26.4	97	35.7	164	3.8	301	24.2
50	23.0	98	3.9	185	60.1	302	3.7
51	3.5	100	44.2	186	12.7	329	12.9
66	17.6	101	6.5	191	7.5	367	7.8
69	82.1	116	4.6	213	39.2	395	7.6
70	15.4	119	100.0	214	7.0	417	8.1
75	29.7	120	26.9	235	9.8	511	5.0
76	4.1	135	62.3	251	25.7	599	4.4

Peaks having intensities lower than 3% of the base peak and lower than m/e 28 are not reported.

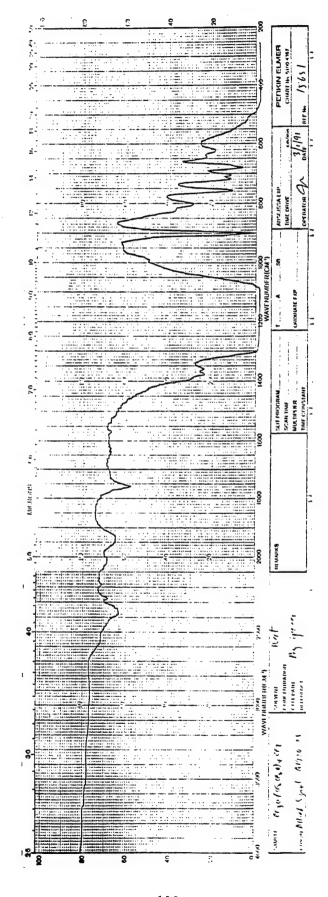
<u>m/e</u>	<u>m/e</u>
599 - [M - F] ⁺	185 - [CF ₃ OCF ₂ CF ₂] ⁺
$417 - [CF_3O(CF_2CF_2O)_2CF_2CF_2]^+$	163 - [FOCCF ₂ CF ₂ O] ⁺
301 - [CF ₃ OCF ₂ CF ₂ OCF ₂ CF ₂] ⁺	$135 - [CF_3OCF_2]^+$
251 - [CF ₃ OCF ₂ CF ₂ OCF ₂] ⁺	$119 - [c_2F_5]^+$
213 - [FOCCF ₂ CF ₂ OCF ₂] ⁺	$69 - [CF_3]^+$

peak. In agreement with the above, the infrared spectrum of the Allied-Signal sample (Figure 58) shows a carbonyl absorption at 1760 cm^{-1} ; the NASA Lewis material does not have this band (see Figure 59).

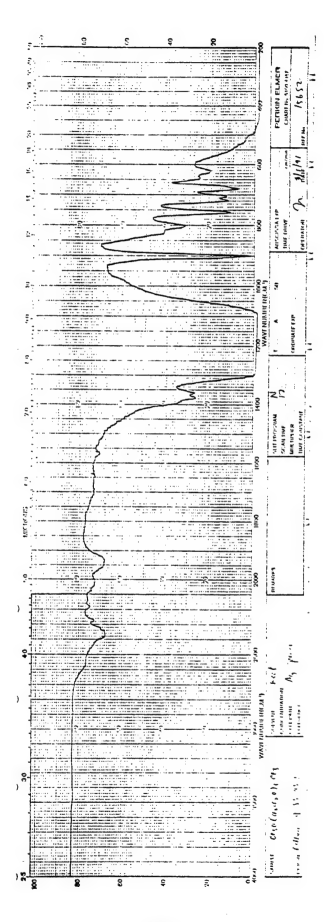
Analysis of the Products Obtained by Allied-Signal from Direct Fluorination of Diglycolic Acid

A 0.95 g sample received from Allied-Signal was subjected to fractionation on the vacuum line through traps kept at 0°C, -78° C, and -196° C. No material was present in the 0°C trap. In the -196° C cooled trap was collected 1.03 mmol of a mixture which consisted of SiF4 (identified by infrared spectroscopy) and a fluorocarbon material which exhibited infrared absorptions at 1918 cm⁻¹ and 1870 cm⁻¹. This is consistent with data reported for perfluoro- β -oxa- δ -valerolactone (bp 30°C, 1866 cm⁻¹ for C=0 stretch [Ref. 25]). The infrared spectrum is shown in Figure 60.

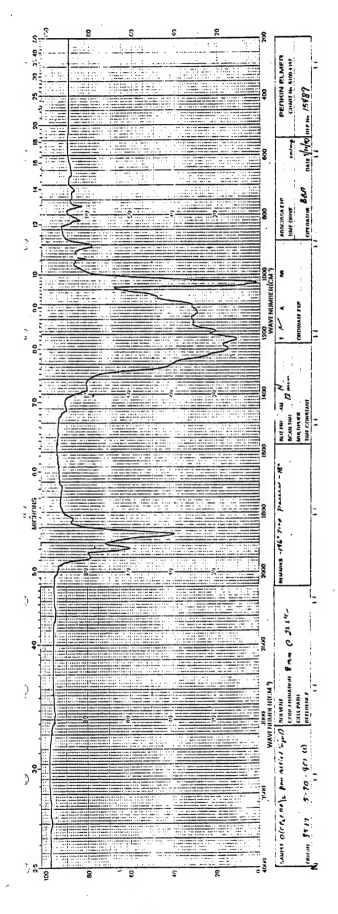
The bulk of the sample (0.78 g) condensed in the -78°C trap. This material exhibited a vapor pressure of 104 mm Hg at 23°C and 43 mm Hg at 0°C. In the infrared spectrum, given in Figure 61, there are several bands in the carbonyl region. None were near 1894 cm⁻¹ reported for O(CF₂COF)₂ [Ref. 25]. The measured vapor pressure does not correspond to the value expected for a material of bp 37-38°C. Although the infrared absorptions in the carbonyl region would indicate the lactone, its bp 29-30°C is even lower than that of the diacetyl fluoride and thus higher vapor pressures should be measured if this was the product.



Infrared spectrum of CF30(CF2CF20)4CF3 prepared by Allied-Signal (0.03 mm Ag spacer). Figure 58.



Infrared spectrum of CF30(CF2CF20)4CF3 received from NASA Lewis (0.03 mm Ag spacer). 59.



-196°C from $O(CF_2COF)_2$ supplied gas phase: 8.0 mm at 23.2°C). Infrared spectrum of the material isolated at by Allied-Signal (IR# 15489, ID# 5-89-40 (1); (IR# 15489, ID# 5-89-40 (1); Figure 60.

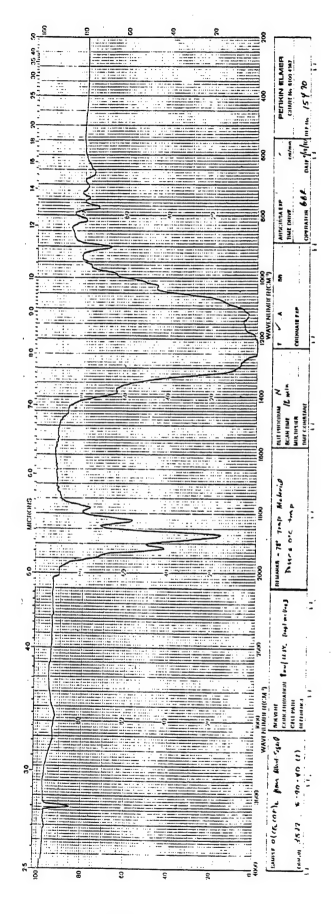


Figure 61. Infrared spectrum of the material isolated at -78°C from O(CF2COF)2 supplied by Allied-Signal (IR# 15490, ID# 5-90-40 (2); gas phase: 8.0 mm at 22.3°C).

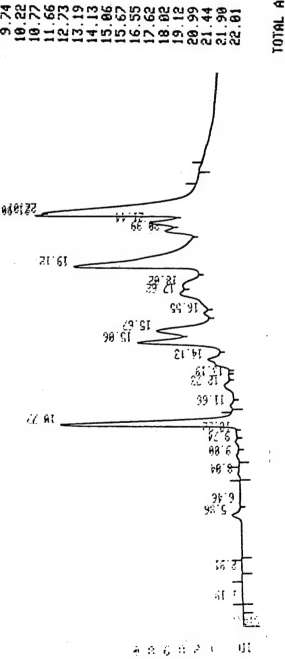
GC analyses, both on the -78°C fraction itself and after the treatment with methanol, are shown in Figures 62 and 63, respectively. The better GC resolution of the methanol-treated material is not surprising in view of the reactivity of the -C(0)Flinkages. The presence of -C(0)F groups was shown by infrared spectroscopy and was confirmed by the formation of methyl esters as determined by GC/MS: m/e 59 $[CO_2Me]^+$ and 109 $[CF_2CO_2Me]^+$. The first peak (Figure 63, 7.77 minutes) was methanol. All the other peaks exhibited the [CO2Me]+ ion indicating the presence of different types of carbonyl fluorides in the original material. Compounds having the arrangements CF2CF2CF2CF2CO2Me (m/e 259; peak at 16.74 minutes) were also present; the GC/MS showed clearly the absence of $O(CF_2CO_2Me)_2$. This means that none of the three precursors discussed above -the acid fluoride, lactone or anhydride- were present in the product mixture received from Allied-Signal. Among the products hydrogenated species were also observed, e.g., in the 15.93 minute GC peak (Figure 63). The m/e 51, [CF2H]+, ion comprised the base (100%) MS peak. Additional evidence of the presence of hydrogenated species were m/e 29 $[CHO]^+$, and/or $[C_2H_5]^+$ and 99 m/e $[C_2H_2F_3O]^+$.

Characterization Data: Experimental fluids MLO 88-48, MLO 88-51, MLO 88-129 and MLO 88-132

 $19_{
m F}$ NMR results for of the fluids listed above are presented in Tables 27-30.

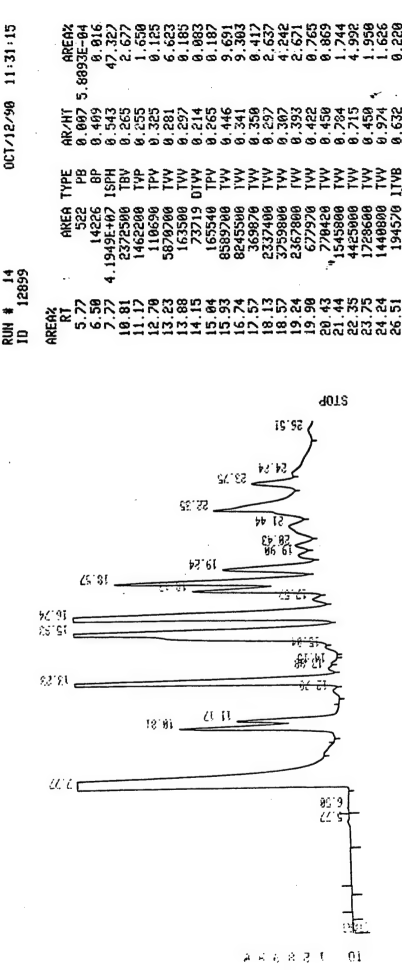
<u> </u>	9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	258882588883 2588855
7876	6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	グイロージョンラフラ
REA 1YPE 163 PB 788 PB 279 PV	746 VB 363 PB 686 PP 739 PP 739 PP 759 PP 768 PP 76	
713/100	21.72 23.23 28.23 161 161 52.53	25.00 23.00 145.00 169.00 12.00 12.00 15.00 15.00 15.00 15.00
AREAZ 1.15 2.9.9	6.46 8.84 9.98 9.74 11.73 14.13	15.55 16.56 16.58 19.59 19.59 19.59

TOTAL AREA= 2.3964E+87 MUL FACTOR= 1.8080E+88;



GC trace of the product mixture (material collected at -78°C, ID# 5-90-40-1) prepared by direct fluorination of diglycolic acid by Allied-Signal. Figure 62.

35-220°C at 8°C/min.; 1.0 µL injected at 200°C, FID at 250°C.) 8' x 1/8" stainless steel packed with Porapak Q, 80/100 mesh. Conditions: (Column:



prepared by direct fluorination of diglycolic acid by Allied-Signal (precursor GC trace of the methanol treated sample (ID# 5-90-40-2) of the material collected at -78°C, ID# 5-90-40-1) Figure 63.

10TAL AREA= 8.8636E+87 MUL FACTOR= 1.0868E+80

35-220°C at 8°C/min.; 1.0 µL injected at 200°C, FID at 250°C.) 8' x 1/8" stainless steel packed with Porapak Q, 80/100 mesh. Conditions: (Column:

TABLE 27

19_F NMR RESULTS FOR MLO 88-48

Sample Identification:

MLO 88-48

Structure Originally Proposed:

 $CF_3O(CF_2CF_2CF_2CF_2O)_nCF_3$

Structure Postulated:

 $\mathtt{CF_3CF_2CF_2O(CF_2CF_2CF_2CF_2O)_nCF_2CF_2CF_3}$

19F NMR Results:

δ ppm	Rel. Int.	Assignment
-80.5	1.0	CF3CF2CF2O-
-128.7	0.7	CF ₃ CF ₂ CF ₂ O-
-82.7	0.7	CF3CF2CF2O-
-81.6	11.8	-OCF2CF2CF2CF2O-
-124.1	11.7	-OCF2CF2CF2CF2O-

Conclusions:

The lack of significant peaks in the region -50~-60 ppm indicates the absence of CF $_3$ O- end group, whereas there is definite evidence for CF $_3$ CF $_2$ CF $_2$ O- arrangement.

TABLE 28

19F NMR RESULTS FOR MLO 88-51

Sample Identification:

MLO 88-51

Structure Originally Proposed:

CF₃O(CF₂CF₂O)_nCF₃

Structure Postulated:

Same as above.

19F NMR Results:

δ ppm	Rel. Int.	Assignment
-54.7	0.9	CF3OCF2CF2O-
-89.4	0.6	CF3OCF2CF2O-
-87.2	14.8	-0C <u>F</u> 2C <u>F</u> 2O-

Conclusions:

The general arrangement is as originally proposed. However, the structure at the base of -89.4 ppm peak would indicate the possibility of a small quantity of material having CF₃CF₂O end groups.

TABLE 29

$19_{ m F}$ NMR RESULTS FOR MLO 88-129

Sample Identification:

MLO 88-129

Structure Originally Proposed:

 $CF_3O(CF_2OCF_2CF_2O)_nCF_3$

Structure Postulated:

Same as above.

19 F NMR Results:

<u>δ ppm</u>	Rel. Int.	Assignment
-56.4	0.8	CF3OCF2O-
-52.7	0.5	CF3OCF2O-
-54.8	0.4	CF3OCF2CF2O-
-89.4	0.3	CF3OCF2CF2O-
-86.2	0.5	CF3CF2OCF2O-
-87.3	0.3	CF3CF2OCF2O-
-50.5	6.7	CF ₃ CF ₂ OC <u>F</u> ₂ O- -OC <u>F</u> ₂ OCF ₂ CF ₂ -
-89.0	14.7	-OCF2OCF2CF2-

Conclusions:

The $^{19}{\rm F}$ NMR results are consistent with the proposed structure. In addition to the end groups CF $_3$ OCF $_2$ O- and CF $_3$ OCF $_2$ CF $_2$ O- originally proposed, CF $_3$ CF $_2$ O- is also present in a significant amount.

TABLE 30

19F NMR RESULTS FOR MLO 88-132

Sample Identification:

MLO 88-132

Structure Originally Proposed:

 $\mathtt{CF_3O[\,(CF_2CF_2O)_4CF_2O]_nCF_3}$

Structure Postulated:

 $CF_3O[(CF_2CF_2O)_3CF_2O]_nCF_3$

19F NMR Results:

δ ppm	Rel. Int.	Assignment
-50.4	1.5	-OCF2OCF2CF2OCF2CF2CF2-
-89.0	3.5	$\hbox{-} \hbox{OCF}_2\hbox{OC}\underline{\hbox{F}}_2\hbox{CF}_2\hbox{OC}\underline{\hbox{F}}_2\hbox{CF}_2\hbox{OC}\underline{\hbox{F}}_2\hbox{-}$
-87.3	7.2	-OCF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ - CF ₃ CF ₂ O-
-54.8	0.7	CF3OCF2CF2O-
-89.4	0.5	CF ₃ OC <u>F</u> 2CF ₂ O-
-56.5	0.1	CF3OCF2O-
-52.7	0.05	CF ₃ OC <u>F</u> 2O-
-86.2	0.1	CF3CF2O-

Conclusions:

If $(CF_2CF_2O)_4$ was present, the ratio of peaks at -89.0 ppm and -87.3 ppm should be 1:3 instead of 1:2; consequently, the arrangement proposed for the major component is $CF_3O[(CF_2CF_2O)_3CF_2O]_nCF_3$. Based on the presence of other low intensity peaks, arrangement CF_3CF_2O - must be also present but in lower concentrations.

For the fluid MLO 88-48 the GC/MS spectra for the three peak pattern (see Figure 1) are given in the order of elution in Tables 31-33.

For the fluid MLO 88-51 the GC/MS spectrum of the two peak pattern (see Figure 2) are given in the order of elution in Tables 34 and 35.

For the fluid MLO 88-129 the representative GC/MS is given in Table 36 and the listing of the prominent ions is compiled in Table 37.

For the fluid MLO 88-132 the representative GC/MS is given in Table 38 and the listing of the prominent ions is compiled in Table 39.

Characterization Data: Chlorofluoropolyalkylether Fluids MLO 90-9 and MLO 88-134

The ¹⁹F NMR data for the fluid MLO 90-9 are listed in Table 40 and the mass spectra of the two peak pattern (see Figure 6a) are given in the order of elution in Tables 41 and 42. The mass spectra of the extra components in the volatiles collected following thermal oxidative degradation of MLO 90-9 (see Figure 8a) are depicted in Table 43.

The 19 F data for the fluid MLO 88-134 are listed in Table 44 and the representative mass spectrum of the two peak pattern (large and small, Figure 7), are compiled in Tables 45 and 46, respectively.

TABLE 31

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF C3F70[CF2CF2CF2CF2C]nC3F7

m/e	%	m/e	%	m/e	%	m/e	%
31	48.5	101	20.0	162	7.8	219	30.2
43	3.5	109	10.2	169	88.6	220	4.8
47	39.9	112	9.2	170	46.1	231	7.7
50	28.6	113	4.5	171	3.4	235	41.5
51	8.4	119	59.7	178	6.7	236	9.0
62	8.8	120	7.0	181	15.9	243	5.1
66	21.2	128	7.6	182	3.0	281	5.3
69	74.9	131	79.6	185	3.8	316	9.7
70	11.0	132	11.3	193	4.8	331	3.6
74	5.9	135	3.3	197	91.3	335	4.0
78	20.1	143	5.2	198	27.3	347	4.9
81	19.9	145	11.5	199	6.3	385	44.4
85	4.9	147	9.5	200	7.5	386	13.7
93	16.6	150	94.9	201	4.5	397	3.5
97	15.9	151	21.8	208	11.5		
100	100.0	159	7.4	216	4.0		

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

m/e	%	m/e	%	m/e	%	m/e	%
31	20.4	100	52.6	159	3.5	219	100.0
47	14.4	101	6.2	169	42.6	220	22.6
50	16.0	109	4.6	170	13.8	235	14.0
51	4.0	112	5.1	181	10.8	247	6.3
62	4.5	119	37.7	185	5.3	269	4.2
66	11.7	120	3.5	197	54.6	285	8.4
69	49.7	131	52.8	198	15.3	316	4.1
70	6.4	132	5.9	199	3.1	335	4.9
78	9.4	145	3.5	200	3.7	347	4.9
81	10.1	150	46.1	201	3.3	385	15.0
93	6.2	151	7.4	208	5.4	435	11.1
97	4.3						

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

TABLE 33

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF $C_2F_5O[CF_2CF_2CF_2CF_2O]_nC_3F_7$

m/e	%	m/e	8	m/e	%	m/e	%
31	27.5	100	63.4	159	4.5	220	6.4
47	22.0	101	9.5	162	3.3	231	5.5
50	21.6	109	6.9	169	41.3	235	16.7
51	6.2	112	5.0	170	25.3	269	12.9
62	5.3	119	100.0	181	18.1	285	6.6
66	21.6	120	11.8	185	22.4	316	6.7
69	64.7	131	53.9	186	3.3	319	9.7
70	7.7	132	5.6	197	81.2	335	22.0
78	11.2	135	4.4	198	18.9	336	5.3
81	11.1	145	4.1	200	3.5	385	13.4
93	9.6	150	77.0	208	8.2	397	3.3
97	5.4	151	12.6	219	44.9		

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

TABLE 34

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF CF30[CF2CF20]nCF3

m/e	%	m/e	8	m/e	%	m/e	%
31	22.4	98	11.0	186	15.7	302	8.0
32	5.3	100	67.9	191	25.5	307	3.0
44	3.5	101	9.1	192	3.9	329	32.0
47	37.4	116	7.2	213	95.3	330	7.9
50	25.0	119	100.0	214	20.8	351	19.2
51	5.1	120	26.8	215	4.0	367	5.2
66	28.9	131	5.9	216	13.0	395	7.1
69	82.8	135	70.8	235	23.9	417	21.2
70	7.5	136	7.7	236	3.1	418	3.0
75	57.2	150	4.5	251	24.7	445	20.4
76	6.5	163	47.8	252	3.3	446	4.3
78	21.0	164	6.6	279	12.1	467	12.7
81	8.7	169	5.8	281	3.3	533	7.9
85	4.0	185	92.9	301	41.4	561	8.7
97	86.6						

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

m/e	%	m/e	%	m/e	8	m/e	%
31	19.2	97	95.3	169	5.9	301	28.8
32	5.9	98	9.2	185	75.5	302	4.8
44	4.0	100	57.5	186	11.1	313	4.1
47	27.3	101	5.6	191	25.8	329	29.3
50	23.1	116	4.9	213	90.7	330	7.0
66	22.5	119	100.0	214	14.2	351	28.7
69	69.1	120	23.2	215	3.6	352	5.2
70	3.9	131	5.2	216	8.6	395	3.3
73	6.0	135	35.7	235	25.6	417	13.1
75	44.2	136	3.2	236	3.5	445	18.3
76	3.9	147	3.2	251	16.2	467	16.0
78	17.2	150	4.0	263	3.0	533	3.3
81	8.2	163	33.9	279	6.2	561	7.7
85	4.5	164	3.7	281	4.6	583	3.9

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

TABLE 36

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF CF₃O[CF₂OCF₂CF₂O]_nCF₃ AND CF₃O[CF₂OCF₂CF₂O]_nCF₂OC₂F₅

m/e	%	m/e	%	m/e	%	m/e	%
31	10.4	100	26.0	191	5.5	302	10.2
32	3.4	101	6.7	197	3.0	329	6.4
44	7.2	113	11.4	201	4.3	341	6.4
47	30.9	116	16.4	213	29.0	342	3.2
50	19.1	119	75.7	214	6.8	351	14.1
51	4.3	120	16.7	235	36.3	355	3.2
66	23.6	135	72.4	236	4.2	367	23.6
69	66.2	136	8.6	251	52.3	395	7.0
75	13.7	147	7.4	252	11.4	417	14.9
78	3.8	163	40.8	263	7.8	429	6.7
81	4.7	164	7.2	279	17.9	430	3.3
85	6.2	169	5.3	281	10.3	433	3.4
97	54.1	185	100.0	282	3.7	483	8.1
98	6.7	186	16.2	301	45.9	511	3.3

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

TABLE 37

LISTING OF PROMINENT IONS SUPPORTING THE STRUCTURE ASSIGNMENT FOR MLO 88-129

CF₃O[CF₂OCF₂CF₂O]_nCF₃ CF₃O(CF₂OCF₂CF₂O)CF₂OC₂F₅ Major Components:

<u>m/e</u>	<u>ion</u>
113	OCF ₂ COF
135	CF ₃ OCF ₂
163	CF2OCF2COF
185	CF ₃ OCF ₂ CF ₂ , C ₂ F ₅ OCF ₂
213	CF2CF2CCF2COF
251	CF3OCF2OCF2CF2, CF3OCF2CF2OCF2
279	CF2OCF2CF2COF
301	C ₂ F ₅ OCF ₂ OCF ₂ CF ₂
317	CF3OCF2OCF2CF2OCF2
351	C ₂ F ₅ OCF ₂ CF ₂ OCF ₂ CF ₂
367	C ₂ F ₅ OCF ₂ OCF ₂ CF ₂ OCF ₂ , CF ₃ OCF ₂ CF ₂ OCF ₂ CF ₂
417	CF ₃ O[CF ₂ CF ₂ OCF ₂ O] ₂
433	$CF_3(OCF_2OCF_2CF_2)_2$, $CF_3(OCF_2CF_2OCF_2)_2$
483	C ₂ F ₅ (OCF ₂ OCF ₂ CF ₂) ₂
549	CF ₃ (OCF ₂ CF ₂ OCF ₂) ₂ OCF ₂ CF ₂ , C ₂ F ₅ (OCF ₂ OCF ₂ CF ₂) ₂ OCF ₂

TABLE 38

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF CF30[(CF2CF20)3CF20]nCF3

m/e	%	m/e	8	m/e	%	m/e	%
31	51.7	109	3.9	186	26.3	330	12.2
32	4.8	112	3.8	187	4.3	351	20.0
40	4.7	113	23.8	191	33.3	352	4.7
44	7.9	114	3.4	192	6.4	367	8.1
47	84.0	116	23.2	194	5.7	395	11.2
48	7.3	117	4.9	197	3.4	417	33.1
50	59.0	119	82.3	213	91.6	418	10.5
51	10.0	120	29.3	214	21.6	445	21.1
66	73.9	128	7.2	215	6.6	446	4.6
67	7.7	131	8.6	216	20.2	467	14.6
69	71.2	135	84.5	217	3.1	468	3.9
70	11.3	136	11.1	235	34.8	483	19.9
75	73.1	137	4.4	236	8.7	484	3.0
76	6.9	144	3.9	251	45.8	511	17.4
78	32.4	147	7.4	252	9.4	512	3.1
79	5.1	150	9.8	259	3.3	533	14.8
81	19.2	159	3.0	260	3.7	561	5.0
85	8.6	163	80.1	263	5.0	582	3.5
87	3.8	164	13.2	279	23.2	599	11.9
93	4.6	166	6.9	280	3.8	627	8.6
94	6.3	167	3.4	282	3.7	649	3.4
97	86.2	169	6.7	301	46.0	715	10.9
98	14.8	181	7.3	302	10.4	765	4.0
100	100.0	185	85.9	329	58.7	809	13.1
101	15.6						

Peaks having intensities lower than 3% of the base peak and lower than m/e 31 are not reported.

TABLE 39

LISTING OF PROMINENT IONS SUPPORTING THE STRUCTURE ASSIGNMENT FOR MLO 88-132

m/e	<u>ion</u>
113	OCF ₂ COF
119	C_2F_5
135	C ₂ F ₅ O, CF ₃ OCF ₂
163	CF2OCF2COF
185	CF3OCF2CF2
213	CF2CF2CCF
235	C ₂ F ₅ OCF ₂ CF ₂
251	CF3OCF2OCF2CF2
301	CF3OCF2CF2CF2
329	CF2CF2OCF2CF2COF
351	C ₂ F ₅ (OCF ₂ CF ₂) ₂
367	CF ₃ OCF ₂ (OCF ₂ CF ₂) ₂
417	CF ₃ (OCF ₂ CF ₂) ₃
445	(CF ₂ CF ₂ O) ₃ CF ₂ COF
483	$CF_3OCF_2(OCF_2CF_2)_3$, $CF_3O(CF_2CF_2O)_3CF_2$
533	C ₂ F ₅ (OCF ₂ CF ₂) ₃ OCF ₂
599	CF ₃ O(CF ₂ CF ₂ O) ₃ CF ₂ OCF ₂ CF ₂
715	CF ₃ O(CF ₂ CF ₂ O) ₃ CF ₂ (OCF ₂ CF ₂) ₂
809	$\mathtt{CF_2O(CF_2CF_2O)_3CF_2OCF_2CF_2OCF_2CF_2OCF_2COF}$
831	$CF_3O(CF_2CF_2O)_3CF_2(OCF_2CF_2)_3$

19_F NMR RESULTS FOR MLO 90-9

TABLE 40

Structure Postulated:

 $ClcF_2CF_2O[CF_2CF(CF_2C1)O]_nX$ $X = CF_3, CF_2CF_2CF_2C1$

19F NMR Data:

mqq	Rel. Int.	<u>Assignment</u>
-52.2	1.1	C <u>F</u> 30-
-73.1	1.5	clcF2cF2o-, clcF2cF2cF2o-
-85.8	1.5	$\operatorname{clcF}_2\operatorname{cF}_2\operatorname{o-}$, $\operatorname{clcF}_2\operatorname{cF}_2\operatorname{cF}_2\operatorname{o-}$
-122.5	0.5	ClcF ₂ CF ₂ CF ₂ O-
-66.0	3.8	-CF ₂ CF(C <u>F</u> ₂ C1)0-
-67.3	0.5	$-CF_2CF(C\underline{F}_2C1)O-$ (end unit)
-75.2 -79.9	2.4 1.5	-C <u>F</u> 2CF(CF ₂ C1)O-
- 77.5	0.5	$-CF_2CF(CF_2C1)O-$ (end unit)
-138~-140	2.5	-CF ₂ C <u>F</u> (CF ₂ C1)0- -CF ₂ C <u>F</u> (CFHC1)0- (?) -CFHC <u>F</u> (CF ₂ C1)0- (?)
-64.8	0.5	?

There is no evidence for the existence of the end group $ClCF_2CF(CF_2C1)O$. The end groups such as $ClCF_2CF_2O$, $ClCF_2CF_2O$, and CF_3O - are present in an approximate ratio of 2:1:1.5.

TABLE 41

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF ${\tt ClCF_2CF_2O[CF(CF_2Cl)CF_2O]_nCF_3}$

m/e	%	m/e	%	m/e	8	m/e	8
31	20.4	96	1.7	150	12.8	251	7.7
35	6.2	97	20.7	151	4.2	253	2.7
36	9.8	99	1.3	153	2.0	263	1.7
37	1.9	100	22.5	163	18.1	279	4.5
38	5.2	101	2.8	164	1.1	281	1.6
39	3.3	109	4.6	165	7.9	282	1.1
47	20.2	112	2.5	166	8.6	295	4.7
50	16.9	113	10.2	167	1.1	297	3.2
51	1.8	115	4.3	168	1.4	301	15.1
62	1.9	116	14.1	169	21.8	302	1.5
66	14.2	118	6.9	170	1.8	303	7.1
67	1.0	119	39.0	184	1.0	313	3.4
68	3.5	120	2.6	185	100.0	317	15.3
69	84.2	128	3.6	186	17.4	318	1.5
70	7.6	131	33.0	187	71.5	319	7.6
75	3.1	132	3.8	188	7.6	329	2.4
78	20.0	134	1.1	197	5.4	345	1.7
81	9.8	135	97.8	201	25.8	347	1.1
85	81.4	136	14.6	202	2.5	395	1.5
86	5.4	137	71.2	203	15.0	411	4.5
87	57.7	138	6.2	213	3.6	413	3.1
88	1.8	144	1.2	229	9.1	499	1.9
93	4.8	147	9.4	231	4.1	501	1.1
94	4.5	149	1.9	·235	4.7		

Peaks having intensities lower than 1% of the base peak and lower than m/e 31 are not reported.

Significant Ions in Support of Structure and Composition

<u>m/e</u>	<u>m/e</u>
499 - $[ClcF_2CF_2[OCF(CF_2C1)CF_2]_2]^+$ $[CF_3O[CF_2CF(CF_2C1)O]_2CF_2]^+$	185 - [ClCF ₂ CF ₂ CF ₂] ⁺
317 - [ClCF ₂ CF ₂ OCF(CF ₂ Cl)CF ₂] ⁺	135 - [ClcF2CF2]+
[CF ₃ OCF ₂ CF(CF ₂ Cl)OCF ₂] ⁺	85 - [ClCF ₂]+
201 - [ClCF ₂ CF ₂ CF ₂ O] ⁺	78 - [CF ₂ CO] ⁺

TABLE 42

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF ${\tt ClCF_2CF_2O[CF(CF_2Cl)CF_2O]_nCF_2CF_2CF_2Cl}$

m/e	%	m/e	%	m/e	%	m/e	%
31	20.1	86	5.6	135	100.0	203	12.6
32	1.9	87	48.8	136	10.5	213	1.2
35	3.4	88	1.5	137	60.3	229	6.8
36	8.5	93	2.7	138	4.0	231	2.0
37	1.1	94	1.8	147	5.5	251	1.4
38	3.4	97	11.4	149	1.0	282	2.2
39	2.3	99	1.2	150	9.4	301	8.2
47	12.6	100	34.2	151	1.4	303	2.8
50	13.9	101	2.9	163	12.2	313	2.8
51	1.3	109	1.5	165	5.2	317	3.8
62	1.3	112	1.3	166	2.1	319	2.7
66	9.7	113	6.3	169	22.3	329	6.3
68	2.4	115	2.1	170	1.3	331	2.2
69	60.8	116	13.7	185	95.7	351	1.8
70	2.2	118	6.4	186	12.0	433	2.9
75	1.9	119	28.8	187	54.6	435	1.8
78	4.8	120	1.8	188	5.6		
81	8.5	131	24.7	201	25.6		
85	92.8	132	1.7	202	1.8		

Peaks having intensities lower than 1% of the base peak and lower than m/e 31 are not reported.

Significant Ions in Support of Structure and Composition

<u>m/e</u>	<u>m/e</u>
433 - [ClCF ₂ CF ₂ CF ₂ CCF ₂ CF(CF ₂ Cl)CF ₂] ⁺	185 - [ClCF ₂ CF ₂ CF ₂] ⁺
317 - [ClCF ₂ CF ₂ OCF(CF ₂ Cl)CF ₂] ⁺	135 - [ClCF ₂ CF ₂] ⁺
251 - [ClCF ₂ CF ₂ CF ₂ OCF ₂] ⁺	85 - [ClCF ₂]+
201 - [ClCF ₂ CF ₂ CF ₂ O] ⁺	

m/e	%	m/e	%	m/e	8	m/e	%
31	12.6	81	10.4	120	1.6	181	2.5
32	2.8	85	35.9	128	1.9	185	6.2
44	1.7	87	17.9	131	100.0	187	1.0
47	6.0	93	7.1	132	9.3	201	2.0
50	7.1	97	8.3	135	59.9	209	1.3
62	1.8	100	23.9	136	1.6	213	1.2
64	2.7	109	33.1	137	28.6	229	1.0
66	4.3	110	1.0	147	4.7	263	1.0
67	2.0	112	4.3	149	1.0	282	25.6
69	28.9	116	4.2	150	45.8	283	1.6
73	1.3	118	1.3	151	1.4	284	9.5
78	1.3	119	44.2	169	1.9		

Peaks having intensities lower than 1% of the base peak and lower than m/e 31 are not reported.

Significant Ions in Support of Structure and Composition

<u>m/e</u>	<u>m/e</u>
282 - [M] ⁺	119 - [C ₂ F ₅] ⁺
150 - [CF ₂ CF ₂] ⁺	109 - [C ₃ F ₃ O] ⁺
135 - [ClCF ₂ CF ₂] ⁺	85 - [ClCF ₂]+
131 - [CF ₂ CF=CF ₂] ⁺	

19_F NMR RESULTS FOR MLO 88-134

TABLE 44

Structure Postulated:

 $\mathtt{ClCF}_2\mathtt{CF}_2\mathtt{O}[\mathtt{CF}_2\mathtt{CF}(\mathtt{CF}_2\mathtt{Cl})\mathtt{O}]_n\mathtt{X}$

 $X = CF_3, CF_2CF_2CF_2C1$

19F NMR Data:

mqq	Rel. Int.	Assignment
-52.1	0.5	C <u>F</u> 30-
-73.0	1.1	clcF2cF2o-, clcF2cF2cF2o-
-85.8	1.1	$\text{clcF}_2\text{c}_{\underline{F}_2}\text{o-}, \text{clcF}_2\text{c}_{\underline{F}_2}\text{c}_{\underline{F}_2}\text{o-}$
-122.4	0.6	ClCF ₂ C <u>F</u> ₂ CF ₂ O-
-65.7	5.8	-CF ₂ CF(C <u>F</u> ₂ Cl)O-
-67.2	0.6	$-CF_2CF(CF_2C1)O-$ (end unit)
-73.0	4.5	OR OR (OR O)
- 79.3	1.2	-C <u>F</u> 2CF(CF2C1)O-
- 77.3	?	$-C_{\underline{F}_2}CF(CF_2C1)O-$ (end unit)
-137.8~-138.7	3.2	-CF ₂ C <u>F</u> (CF ₂ C1)0-
-64.7	0.5	
-78.7	1.3	
-143.0	0.4	?
-139.5	0.2	

The end groups $ClCF_2CF_2O$ -, $ClCF_2CF_2CF_2O$ -, and CF_3O - are present in an approximate ratio of 1.5:1.8:1.0. The NMR results indicate that this sample should have a higher molecular weight than MLO 90-9.

TABLE 45

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF

m/e	%	m/e	%	m/e	%	m/e	%
31	15.7	93	3.5	149	2.6	279	1.6
32	2.6	94	2.9	150	13.9	285	1.3
35	4.6	96	1.1	151	4.3	301	12.8
36	15.1	97	14.9	153	1.9	302	1.0
37	1.5	100	22.2	163	22.6	303	4.9
38	6.2	101	2.2	164	2.1	313	2.2
39	3.2	109	3.9	165	12.1	317	11.1
40	1.0	112	2.2	166	7.5	319	8.8
47	15.5	113	8.8	167	1.6	321	1.3
50	11.8	115	3.7	168	3.2	329	5.8
51	2.0	116	12.5	169	17.3	331	2.1
62	1.3	118	4.8	170	1.8	345	2.8
66	11.0	119	20.6	185	100.0	347	1.8
68	1.7	120	1.4	186	22.3	351	3.2
69	60.4	128	3.0	187	78.8	367	3.6
70	2.8	131	41.6	188	8.9	369	2.4
78	3.7	132	4.1	201	17.1	433	1.0
81	8.3	135	92.4	202	1.0	499	1.3
85	82.7	136	11.4	203	9.6	549	0.8
86	4.3	137	71.6	213	1.9	681	0.1
87	38.1	138	5.2	.229	4.3	731	0.1
88	1.6	147	12.4	231	1.8		

Peaks having intensities lower than 1% of the base peak and lower than m/e 31 are not reported. High mass peaks which are less than 1% but which are important to structure determination are also listed.

Significant Ions in Support of Structure and Composition

m/e	
	$[ClcF_2CF_2CF_2[OCF(CF_2C1)CF_2]_3]^+$
681 -	[ClCF ₂ CF ₂ [OCF ₂ CF(CF ₂ Cl)] ₃] ⁺
	$[ClcF_2CF_2CF_2[OCF(CF_2Cl)CF_2]_2]^+$
	[ClCF ₂ CF ₂ [OCF ₂ CF(CF ₂ Cl)] ₂] ⁺
	$[CF_3[OCF(CF_2C1)CF_2]_2]^+$
	[ClCF ₂ CF ₂ CF ₂ OCF(CF ₂ Cl)CF ₂] ⁺
	[ClCF ₂ CF ₂ OCF ₂ CF(CF ₂ Cl)] ⁺
	[ClcF ₂ CF ₂ CF ₂ O] ⁺ , [ClcF ₂ CF ₂ OCF ₂] ⁺
	[ClCF ₂ CF ₂ CF ₂] ⁺
135 -	[ClCF ₂ CF ₂] ⁺

TABLE 46

ION FRAGMENTS AND INTENSITIES RELATIVE TO BASE PEAK OF ClCF₂CF₂O[CF₂CF(CF₂Cl)O]_nCF₂CF(CF₃)OCF₂CF₂Cl

m/e	%	m/e	%	m/e	%	m/e	%
31	12.7	101	1.4	165	7.0	301	19.0
32	4.8	109	1.7	166	5.5	302	1.7
35	2.3	112	1.3	168	2.1	303	8.2
36	9.6	113	11.0	169	43.4	313	3.4
38	4.2	115	3.9	170	3.4	317	6.0
39	1.3	116	8.8	185	100.0	319	4.4
47	11.2	118	2.8	186	13.5	329	9.7
50	8.0	119	32.2	187	48.0	331	3.0
66	5.7	120	1.4	188	5.1	335	1.5
68	1.0	128	1.5	197	1.0	345	1.2
69	35.4	131	31.9	201	13.7	351	27.8
70	1.6	132	2.6	203	8.2	352	3.2
78	2.4	135	96.7	207	1.7	353	12.9
81	4.0	136	7.6	213	5.3	367	1.7
85	55.9	137	41.6	229	2.3	483	1.1
86	2.3	138	2.8	235	1.8	499	0.4
87	34.3	147	10.3	251	1.0	533	1.2
93	1.8	149	1.9	263	1.1	715	0.2
94	1.5	150	19.1	279	1.4		
97	19.3	151	2.8	285	4.4		
100	23.0	163	19.3	297	1.2		

Peaks having intensities lower than 1% of the base peak and lower than m/e 31 are not reported. High mass peaks which are less than 1% but which are important to structure determination are also listed.

Significant Ions in Support of Structure and Composition

m/e	
715 -	- [ClCF ₂ CF ₂ CF ₂ OCF(CF3)CF ₂ [OCF(CF ₂ Cl)CF ₂] ₃] ⁺
533 -	- [ClcF2CF2CF2OCF(CF3)CF2[OCF(CF2C1)CF2]2]+
	- $[ClcF_2CF_2[OCF_2CF(CF_2C1)]_2]^+$
351 -	- [ClCF ₂ CF ₂ CF ₂ OCF(CF ₃)CF ₂]+
	- [ClCF ₂ CF ₂ OCF ₂ CF(CF ₂ Cl)] ⁺
	- [ClCF ₂ CF ₂ CF ₂] ⁺
	- [ClCF ₂ CF ₂]+
	- [clcF ₂] ⁺

NMR Data: Commercial Fluids Krytox 143AC, Demnum S-20, Demnum S-100 and Aflunox 2507

Proton and 19 F NMR spectra of Krytox 143AC (MLO 71-6) were reported [Ref. 26]. The multiscan 1 H and 19 F NMR spectra of the fluorinated Krytox 143AC (MLO 91-21) are presented in Figures 64 and 65, respectively. The corresponding NMR spectra of Aflunox 2507 (MLO 88-298) are given in Figures 66 and 67.

The $^{1}\mathrm{H}$ and $^{19}\mathrm{F}$ NMR spectra of Demnum S-20 (MLO 88-177) are depicted in Figures 68 and 69. The chemical shift assignments are listed in Table 47. The corresponding data for Demnum S-100 are presented in Figures 70 and 71 and Table 48.

19F NMR Data: Second Series of Experimental Fluids

The 19 F NMR shifts together with intensities and assignments are presented for the fluids MLO 91-87, 88, 105, 106, 126, 127, 132, 157, 158, 160 and 161 in Tables 49-59, respectively.

Fractional Distillation Data of Perfluoropolyalkylether Fluids

The fractional distillation data for selected Experimental fluids are listed in Table 60.

Thermal Oxidative Stability Testing of Perfluoropolyalkylether Fluids Summary of Results

All the thermal oxidative testing details for the commercial and second series experimental fluids are compiled in Table 61.

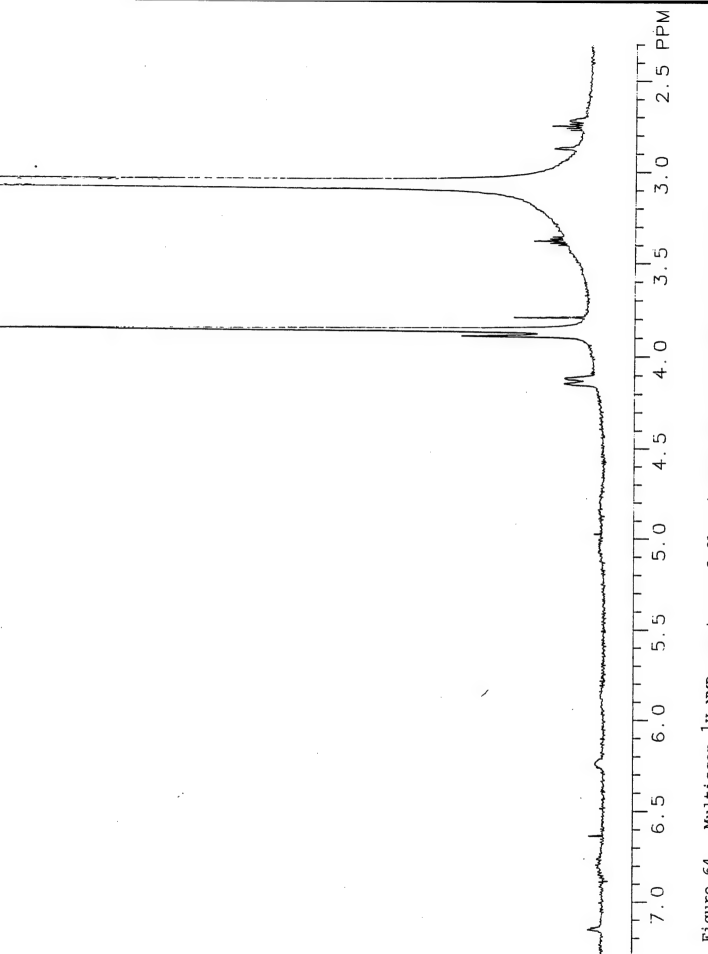
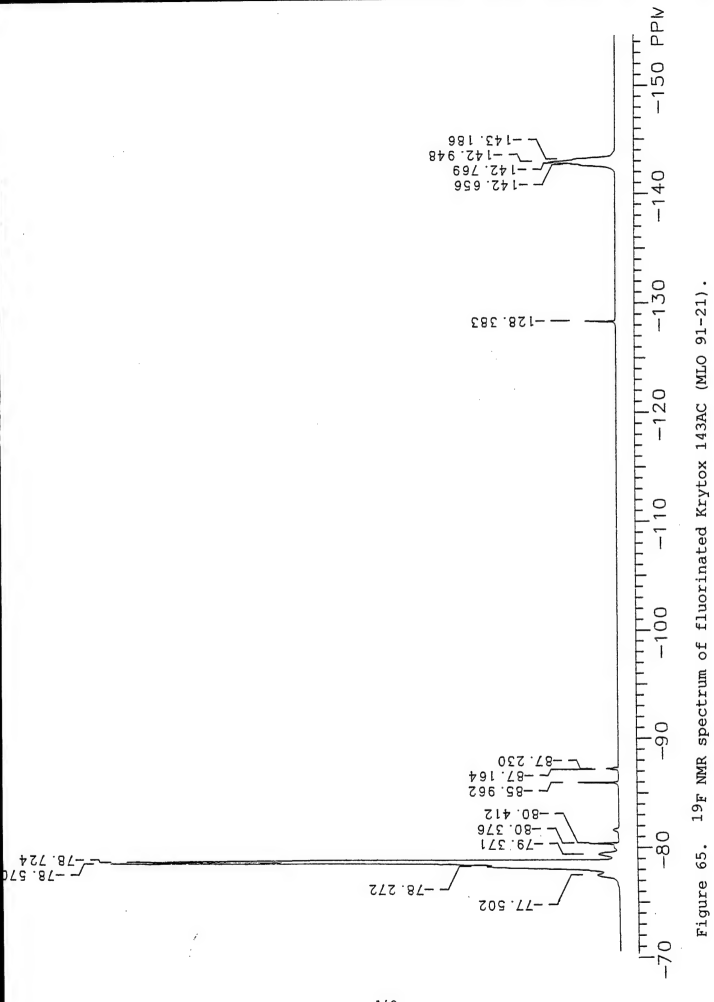
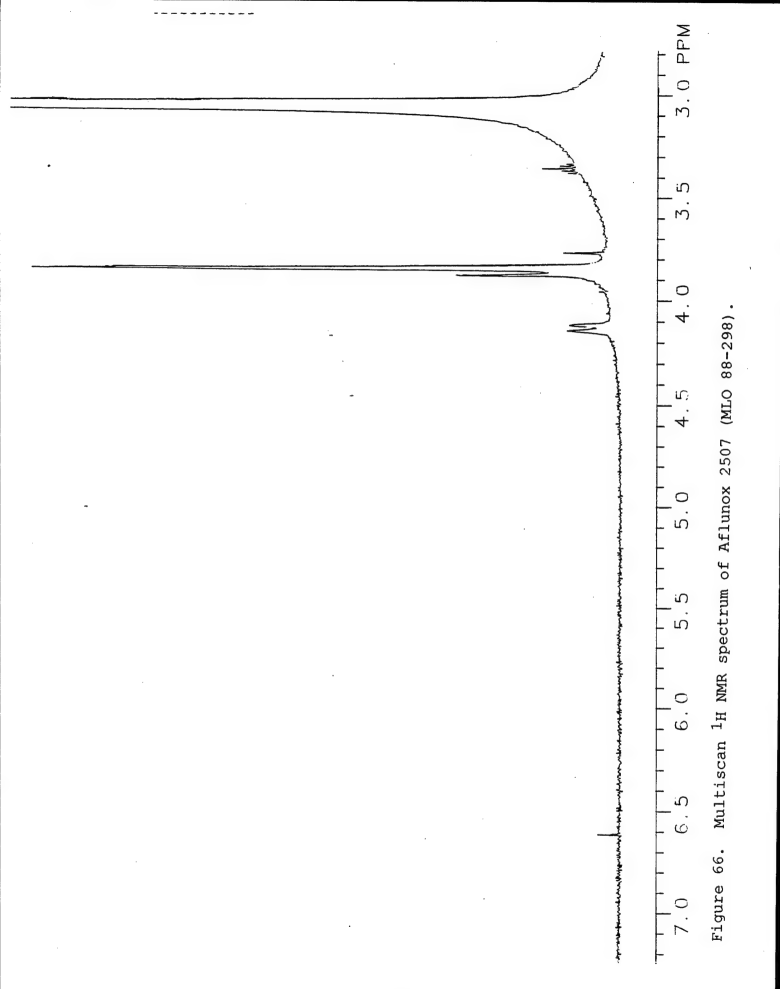
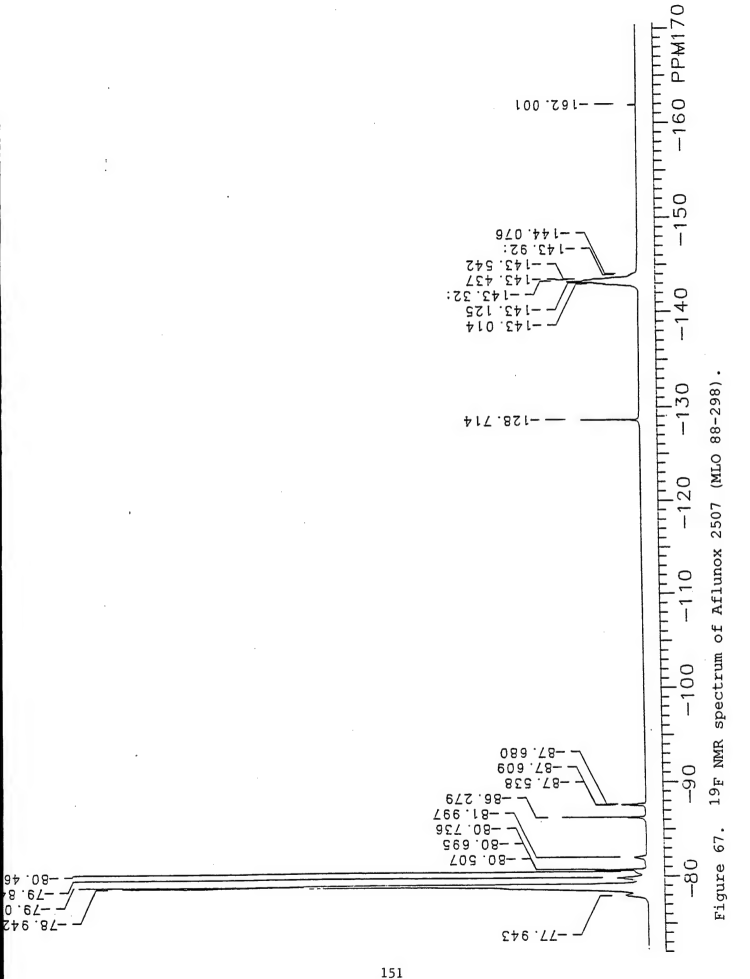


Figure 64. Multiscan ¹H NMR spectrum of fluorinated Krytox 143AC (MLO 91-21).







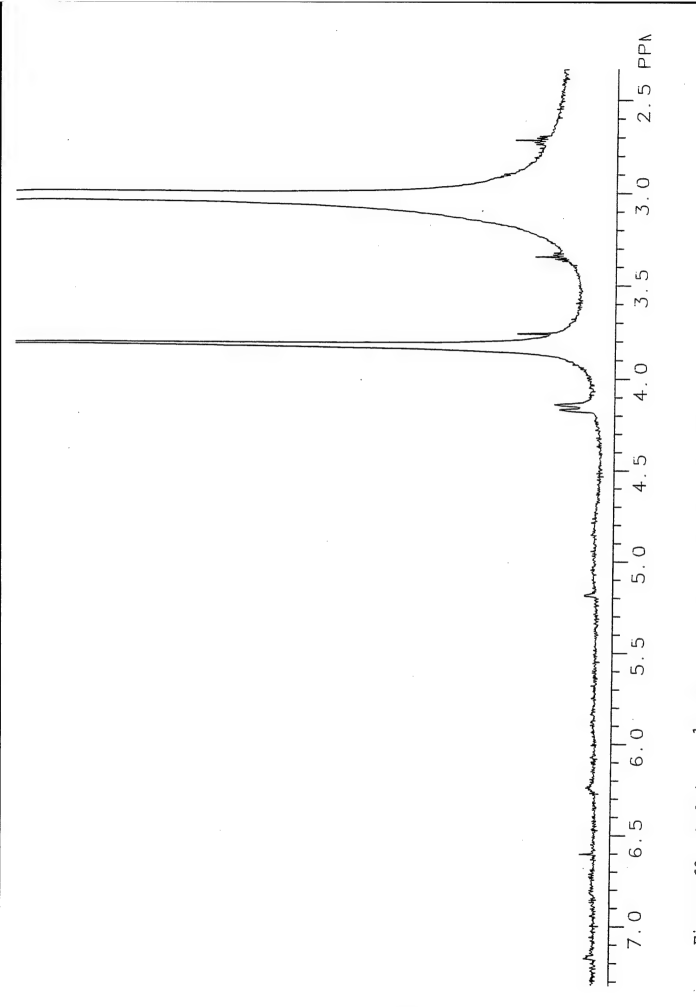


Figure 68. Multiscan $^{1}\mathrm{H}$ NMR spectrum of Demnum S-20 (MLO 88-177).

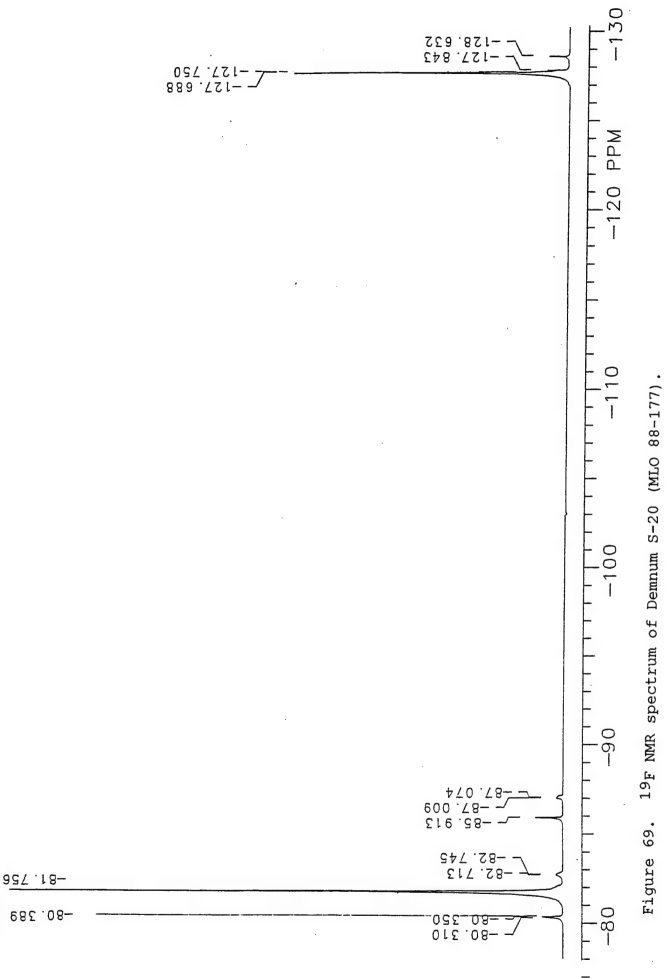
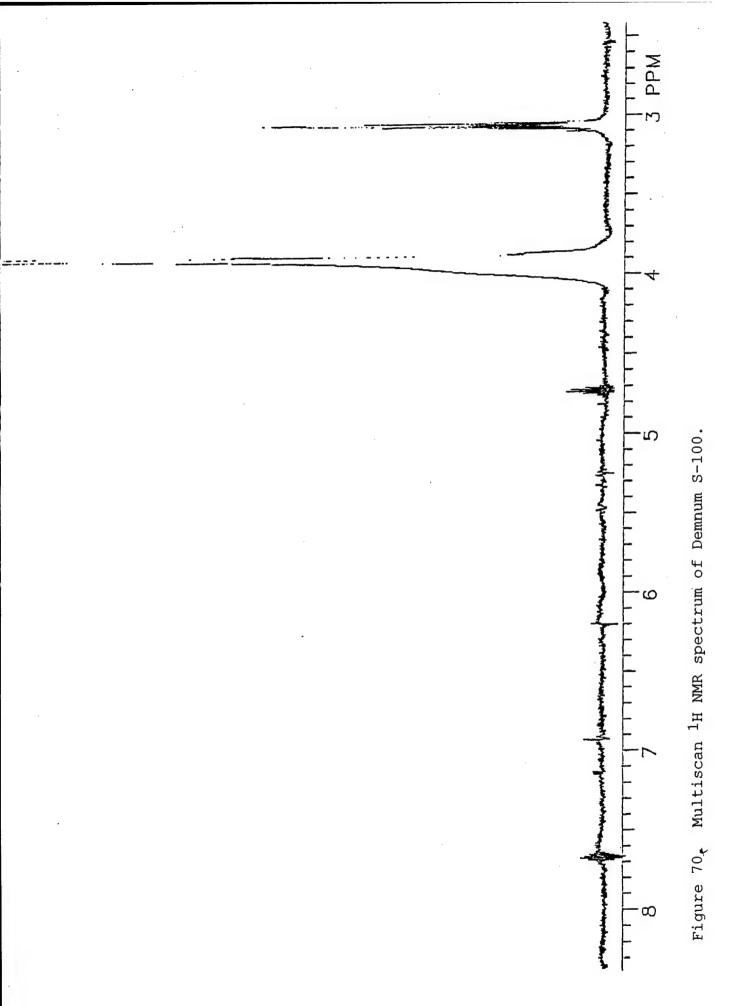


TABLE 47

19F NMR CHEMICAL SHIFT ASSIGNMENT OF DEMNUM S-20 (MLO 88-177)

(ppm)	Rel. Int.	Assignment
-80.4	0.4	CF3CF2CF2O-
-81.8	7.9	-0C <u>F</u> 2CF2C <u>F</u> 20-
-82.7	0.3	CF3CF2C <u>F2</u> O-
-85.9	0.4	CF3CF20-
-87.0	0.25	CF3C <u>F2</u> 0-
-127.7	3.9	-ocf ₂ c _{f2} cf ₂ o-
-128.6	0.25	CF3CF2CF2O-



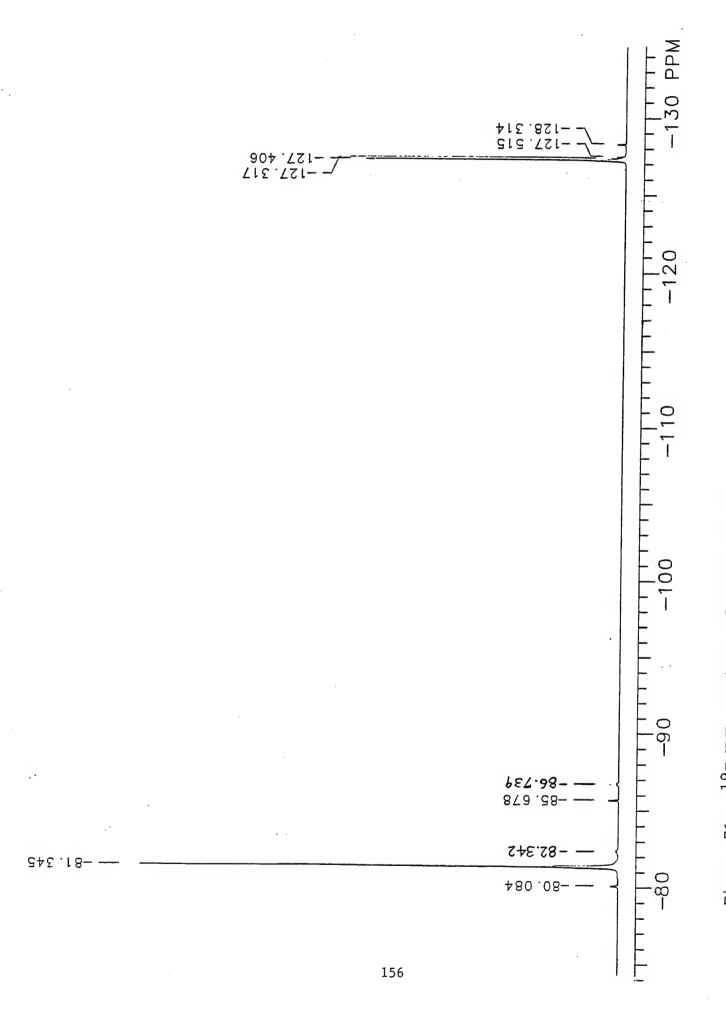


Figure 71. 19F NMR spectrum of Demnum S-100.

TABLE 48 $$^{19}{
m F}$ NMR CHEMICAL SHIFT ASSIGNMENT OF DEMNUM S-100

(ppm)	Rel. Int.	Assignment
-80.1	0.27	CF3CF2CF2O-
-81.3	13.1	-0C <u>F</u> 2CF2C <u>F2</u> 0-
-82.3	0.2 (?)	CF3CF2CF2O-
-85.7	0.27	CF3CF2O-
-86.7	0.17	CF3C <u>F2</u> 0-
-127.4	6.5	-ocf ₂ cf ₂ cf ₂ o-
-128.3	0.18	CF3CF2CF2O-

End group ratio of $CF_3CF_2CF_2O-: CF_3CF_2O-=1:1$ Calculated MW = 6300

TABLE 49 $^{19}{\rm F~NMR~of-O(CF_2CF_2O)_m[CF_2CF(CF_3)O]_n(CF_2CF_2O)_p-~(MLO~91-87)^a}$

mqq	Rel. Int.	Assignment	
-52.7 -54.8	0.1	C <u>F</u> 30-	
-77.3(shou	lder) lder) 3.0	-C <u>F</u> 2CF(C <u>F</u> 3)0-	
-78.6(main	peak)	_2 (_3)	
- 80.5	0.05	CF3CF2CF2O-	
-81.0	0.25	$(C\underline{F}_3)_2CFO-$	
-84.3	0.05	?	
-86.1	b	CF ₃ CF ₂ C <u>F</u> ₂ O-	
-87.3	0.3	-oc <u>F</u> 2c <u>F</u> 2o-	
-89.4	b	CF3OCF2CF2O-	
-128.3	b	CF ₃ C _{F₂} CF ₂ O-	
-138.5	b	(CF ₃) ₂ C <u>F</u> 0-	
-142.9	0.6	-CF ₂ C <u>F</u> (CF ₃)O-	
-144.3	b	?	

a Based on the data listed below the ratio of $CF_2CF(CF_3)$ 0 to CF_2CF_2O is approximately 8:1. The end groups CF_3 , $(CF_3)_2CF$ and $CF_3CF_2CF_2O$ are present in an approximate ratio of 4:2.5:1.

b The relative intensity < 0.05.

TABLE 50

ppm	Rel. Int.	Assignment
-52.7 -54.8	0.1	CF3OCF(CF3)CF2O- CF3OCF2CF2O-
- 78.9	1.7	$-c_{\underline{F}_2}c_{\underline{F}}(c_{\underline{F}_3})o_{\underline{-}}$
-80.5	0.1	$CF_3CF_2CF_2O-$?
- 81.5	0.5	-OCF2CF (OR) CF2O
-84.4	0.5	-ocf ₂ cf(cf ₃)ocf ₂ cf ₂ o-
-87.2	2.4	-oc <u>F</u> 2c <u>F</u> 2o-
-89.2	0.1	CF3OCF2CF2O-
-143.4	0.4	-CF ₂ C <u>F</u> (CF ₃)O-, -CF ₂ -C <u>F</u> -CF ₂ -

The ratio of CF_2CF_2O to $CF_2CF(CF_3)O$ is approximately 1.8:1. The end groups are $CF_3OCF(CF_3)CF_2O$, $CF_3CF_2CF_2$ — and $CF_3OCF_2CF_2$ in the ratio of 1:1:1.5.

TABLE 51 $$^{19}{\rm F}$ NMR of XO(CF2CF2CF2CF2CF2OCF2O) $_{\rm n}{\rm X},~$ MLO 91-105

pm	Rel. Int.	Assignment
-49.9	3.5	-oc <u>F</u> 20-
-52.0	0.15(?)	-0 <u>F</u> 20C <u>F</u> 20-
-54.5	0.4	$CF_3O(CF_2)_4O-$
-56.4	ā	C <u>F</u> 30CF20-
-80.6 C <u>F</u> 3CF ₂ CF	0.8 ₂ CF ₂ O-	$CF_3CF_2CF_2O-$ (major),
-81.7	a	-CF ₂ CF ₂ OC <u>F₂</u> CF ₂ CF ₂ CF ₂ O-
-83.5	6.8	-oc <u>F</u> 2cF2cF2cF2ocF2o-
-84.7	0.5	CF ₃ CF ₂ C <u>F</u> ₂ O-
-86.1	a	CF3CF2O-
-87.2	a	$CF_3C\underline{F}_2O$ -, $-OC\underline{F}_2C\underline{F}_2O$ -
-124.2	7.0	-ocf ₂ c <u>f</u> 2cf ₂ cf-
-124.9	b	CF ₃ CF ₂ CF ₂ CF ₂ O-
-128.8	0.5	CF ₃ C <u>F₂</u> CF ₂ O-
-130.0	a	?

a Peak is of extremely low intensity.

b Peak intensity cannot be determined due to poor resolution.

Table 52 $$^{19}{\rm F}$$ NMR of XO(CF2CF2CF2CF2CF2OCF2O) $_{\rm n}{\rm X}$ MLO (91-106)

ppm	Rel. Int.	Assignment
-49.8	3.5	-oc <u>F</u> 20-
-52.0	a .	-oc <u>F</u> 2oc <u>F</u> 2o-
-54.5	0.4	C <u>F</u> 30-
-56.5	a	CF3OCF2O-
- 76.7	a	?
-80.4	0.9	CF3CF2CF2CF2O-
-81.2	а	?
-81.5	a	?
-83.3(m	•	-cF2cF2cF2cF2ocF2o-
-83.8(s	6.3 houlder)	CF ₃ CF ₂ CF ₂ C <u>F</u> ₂ O-
-84.7	a	$CF_3CF_2C\underline{F}_2O$
-86.2	b	CF3CF2O-
-89.0	b	-ocf ₂ c <u>f</u> 2ocf ₂ o-
-119.1	a	?
-120.6	2.9	-CF ₂ CF ₂ CF ₂ CF ₂ CF ₂ OCF ₂ O-
-121.4	b	?
-124.1	5.9	-CF ₂ CF ₂ CF ₂ CF ₂ CF ₂ OCF ₂ O-
-125.1	1.1	CF ₃ C <u>F</u> ₂ CF ₂ CF ₂ O-
-128.8	b	CF ₃ C <u>F</u> ₂ CF ₂ O-
	_	

a Peak is of extremely low intensity.

b Peak intensity cannot be determined due to the poor resolution or low intensity.

TABLE 53 $$^{19}{\rm F}$ NMR of X(CF2CF2CF2CF2C) $_{\rm n}{\rm X}$ (MLO 91-126)

		•
ppm	Rel.Int.	Assignment
-54.6	a	CF3OCF2CF2CF2CF2O-
-80.5	0.9	$CF_3CF_2CF_2O$ - (major), $CF_3CF_2CF_2CF_2O$ -
-81.6	7.6	-c <u>F</u> 2cF2cF2c <u>F</u> 2o-
-82.7	0.6	$CF_3CF_2C\underline{F}_2O$ - (major), $CF_3CF_2CF_2C\underline{F}_2O$ -
-83.9	a	?
-84.2	a	?
-86.1	b	CF3CF2O-
-87.2	a	CF ₃ C <u>F</u> ₂ 0-
-109	a	?
-121.3	a	?
-122.0	a	?
-124.0	7.6	-CF ₂ C <u>F</u> ₂ CF ₂ CF ₂ O-
-125.0	b	CF ₃ C <u>F₂CF₂CF₂C</u> -
-128.3	a	?
-128.7	0.6	CF ₃ C <u>F₂</u> CF ₂ O-
-129.1	a	?
-130.7	a	?
-131.5	a	?

a Peak is of extremely low intensity.

b Peak intensity cannot be determined due to the poor resolution or low intensity.

TABLE 54 19 F NMR OF $_{\rm XO[CF_2CF(CF_3)O]_nX}$ (MLO 91-127)

ppm	Rel.Int.	Assignment
-52.7		$C_{\underline{F}_3}$ OCF(CF ₃)CF ₂ O-
-54.5		$C_{\underline{F_3}}OCF_2CF(CF_3)O$
- 56.0	0.01	CF3OCF2CF2O-
-59.9	a	?
-70.0	b(broad)	?
-72.2	b	ClCF2CF2CF2O-
-73.1	a	C1C <u>F</u> 2CF ₂ O-
-77.3	7	- ·
-78.1	- 6.7	
-78.6	major	$-\text{OC}\underline{F}_2\text{CF}(\text{C}\underline{F}_3)\text{O}-$
- 79.6	0.1	?
-80.5	0.15	CF3CF2CF2O-
-81.1	0.45	$(C\underline{F}_3)_2CFO-$
-82.8	b	-OC <u>F</u> 2CF2C <u>F</u> 2O-
-83.4	b	-OCF2CF2CF2CF2O
-84.1	b	?
-86.1	0.09	C <u>F</u> 3CF ₂ O-
-87.1	0.06	$CF_3C\underline{F}_2O-$, $-OC\underline{F}_2C\underline{F}_2O-$?
-89.0	a	$CF_3OC\overline{F}_2CF_2O-?$
-120.9	b	?
-121.2	b	?
-124.0	b	-ocf ₂ c <u>f</u> 2cf ₂ cf-
-125.9	b	?
-127.2	b	-OCF ₂ CF ₂ CF ₂ O-
-128.4	major 0.1	$CF_3C\overline{F}_2C\overline{F}_2O\overline{-}$
-128.5	J	
-138.2	0.08(broad)	(CF ₃) ₂ C <u>F</u> O-
-142.6	1.3	$-\text{OCF}_2\overline{\text{C}}_{\underline{\text{F}}}(\text{CF}_3)$ O-
-144.3	0.05	$CF_3OC_F(CF_3)CF_2O-$

a Peak is of extremely low intensity.b Peak intensity cannot be determined due to poor resolution or low intensity.

TABLE 55 19 F NMR OF $XO[CF_2CF(CF_3)OCF_2CF(CF_3)OCF_2O]_nX$ (MLO 91-132)

mqq	Rel.Int.	Assignment	
-44.5 -45.1	a	-OCF ₂ CF(CF ₂ CF ₃)OC <u>F₂OCF(CF₂CF₃)CF₂O-</u>	
-45.1 -45.6	a 0 4	(OCE OF OCE	
-47.7	0.4 0.7	-OCF ₂ CF(CF ₃)OC _{F2} OCF(CF ₃)CF ₂ O-	
	major ₁	-OCF ₂ CF(CF ₃)OCF ₂ OCF ₂ CF(CF ₃)O-	
43.3	- 0.3	$-\text{OCF}(\text{CF}_3)\text{CF}_2\text{OCF}_2\text{OCF}_2\text{CF}(\text{CF}_3)\text{O-}$	
-50.1] "."	-OCF ₂ CF ₂ OCF ₂ OCF ₂ CF(CF ₃)O-	
-52.7	0.13	$C\underline{F}_3$ OCF(CF ₃)CF ₂ O-, -OC \underline{F}_2 OCF ₂ O-?	
-54.7	0.2	CE_3 OCF ₂ CF(CF ₃)O-	
-56.5	b	CF30CF2CF20-	
	shoulder?	- <u>-</u> J 2 2	
-78.5	4.5	$-OC_{\underline{F}_2}CF(C_{\underline{F}_3})O-$	
-79.3	0.3	$-0C\underline{F}_2CF(CF_2C\underline{F}_3)O-$	
-80.5	0.12	$C_{\underline{F}_3}C_{\underline{F}_2}C_{\underline{F}_2}C_{\underline{F}_3}$	
-81.3	0.9	$(\overline{CF_3})_2^{\circ} CFO-$	
-82.8	1.2	?	
-84.6	b	-oc <u>F</u> 2cF2cF2c <u>F</u> 2o- ?	
-86.1	0.1	$C\underline{F}_3CF_20-$	
-87.3	0.5	-OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ O- major, C	Fac
-89.0	0.15	-OCF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ O-	-
- 89.5	0.06	$CF_3OC\underline{F}_2CF_2O$	
-122.6	l h	OCE CE/CE CE \o	
-122.7	} b	$-\text{OCF}_2\text{CF}(\text{C}_2\text{CF}_3)\text{O}-$	
-124.0	b	-OCE-CE-CE-CE-O-	
-125.0	a	-0CF ₂ C <u>F₂CF₂CF₂O-</u> CF ₃ C <u>F₂CF₂CF₂O-</u>	
-127.6	a	CF ₃ CF ₂ CF ₂ OCF ₂ OCF(CF ₃)CF ₂ O-?	
-127.7	a	CF ₃ C _E ₂ CF ₂ OCF ₂ OCF ₂ CF(CF ₃)O-?	
-128.5	1	CF ₃ C _{E₂} CF ₂ OCF (CF ₃) CF ₂ O-	
	- 0.09	2 -= 7 21 7 22 (21 3) 21 50 -	
-128.7]	CF ₃ C <u>F₂</u> CF ₂ OCF ₂ CF(CF ₃)O-	
-138.7	0.15	(CF ₃) ₂ C <u>F</u> O-	
-141.2	a	-OCF ₂ CF ₂ OCF ₂ C <u>F</u> (CF ₂ CF ₃)OCF ₂ O-	
-142.0	a	-OCF ₂ OCF ₃ CF(CF ₂ CF ₃)OCF ₂ CF ₂ O-	
-143.5	0.3	$-CF_2C\underline{F}(CF_3)OCF_2OC\underline{F}(CF_3)CF_2O-$	
-144.6	0.8	-OCF ₂ OCF ₂ CF(CF ₃)OCF ₂ CF(CF ₃)O-	

a Peak is of extremely low intensity.b Peak intensity cannot be determined due to poor resolution or low intensity.

TABLE 56 $$^{19}{\rm F}$ NMR of XO[(CF2CF2O)4CF2O]nX (MLO 91-157)

ppm	Rel.Int.	Assignment
-47.4		-CF ₃ CF ₂ OC <u>F</u> ₂ O- ?
-49.5	a	$-\text{OCF}(\text{CF}_3) \overline{\text{OC}}_{\text{2}} \text{OCF}_2 \text{CF}_2 - ?$
-50.4	0.8	$-(CF_2CF_2O)_4C\overline{F}_2O(CF_2CF_2O)_4-$
-52.7	b	$CF_3OC_{F_2}O$
-54.6	0.6	$CF_3(OCF_2CF_2)_XO-$
-54.8 (majo	or)	_5
-56.5	a	C <u>F</u> 3OCF2O-
-73.0	a	CICF2CF2O ?
-78.4	a	$-0C\underline{F}_2CF(C\underline{F}_3)O-?$
-79.2	a	side chain CF3
-80.5	a	$-c_{\underline{F}_3}c_{\underline{F}_2}c_{\underline{F}_2}o_{\underline{F}_3}$
-81.6	a	$(\overline{C}\underline{F}_3)$ 2CFÖ- ?
-83.1	a	?
-83.9	a	?
-85.1	a	$-OF(CF_3)-?$
-86.2	6.2	C <u>F</u> 3CF20-
-87.3	5.5	-OCF2OCF2CF2O(CF2CF2O)2CF2CF2O-
-89.0	1.7	$-\text{OCF}_2\text{OC}_2\text{CF}_2\text{CF}_2\text{O}$ ($\text{CF}_2\text{CF}_2\text{O}$) $_2\text{CF}_2\text{C}_2\text{C}_2\text{O}$
-109.6	a	?
-122.7	a	$-\text{OCF}_2\text{CF}(\text{C}_2\text{CF}_3)\text{O}$?
-124.0	a	$-\text{OCF}_2\text{C}_2\text{C}_2\text{C}_2\text{C}_2\text{C}_2$

a Peak is of extremely low intensity

b Peak intensity cannot be determined due to the poor resolution or low intensity.

TABLE 57 19F NMR OF XO[CF₂CF(CF₂OCF₂CF₂OCF₂CF₂OCF₃)0]X (MLO 91-158)

mqq	Rel.Int.	Assignment
-50.4	a	-oc <u>F</u> 20-
- 52.3	0.15	$C\underline{F}_3$ OCF(R)CF ₂ O-, $-OC\underline{F}_2$ OC \underline{F}_2 O- ?
-54.4] 1.8	
-54.7	major	-CF ₂ CF(CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OC <u>F₃</u>)O-
-55.5	b	?
- 76.6	0.9	-CF2CF(CF2OCF2CF2OCF2CF3)O-
-78.5	b	$-0\overline{C}_{2}^{F}CF(C\overline{E}_{3})0-?$
-80.2	1.1	-CF ₂ CF(CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₃)O-
-81.9	b	$-\text{OC}_{\underline{F}_2}\text{CF}_2\text{CF}_2\text{CF}_2\text{O}$, $-\text{OC}_{\underline{F}_2}\text{CF}_2\text{CF}_2\text{O}$
-83.6	b(broad)	?
-85.1	b(broad)	$-OCF(C\underline{F}_3)O-?$
-86.1	b	CF_3CF_2O
- 87.3	3.5	-CF ₂ CF(CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₃)O- (major), CF ₃ CF ₂ C
-89.3	1.2	-CF ₂ CF(CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₃)O-
-124.4	b	$-\text{OCF}_2\text{C}_2\text{C}_2\text{C}_2\text{C}_2\text{O}$
-127.2	b	-OCF ₂ CF ₂ CF ₂ O-
-142.0	ו ו	·
		$-OCF_2C\underline{F}(R)O-$
-142.6	- 0.6 J	
-144.0	J	$CF_3OC\underline{F}(R)CF_2O-$

a Peak is of extremely low intensity.
b Peak intensity cannot be determined due to poor resolution or low intensity.

TABLE 58 $$^{19}{\rm F}$ NMR of XO(CF2CF2OCF2O) $_{\rm n}{\rm X}$, MLO 91-160

ppm	Rel. Int.	Assignment
-47.3	b	-ocf(cF ₃)oc <u>F₂ocF₂cF₂o-</u> ?
-50.5	2.3	-ocf ₂ cf ₂ oc _{f2} cf ₂ cf ₂ o-
-51.4	a	?
-52.0	a	-oc <u>f</u> 2oc <u>f</u> 2o-
-52.8	0.2	CF ₃ OC <u>F</u> ₂ O-
-54.8	0.17	CF3OCF2CF2O-
- 55.9	a	CF_3 OCF(CF_3)O-?
-56.5	0.3	c <u>F</u> 30cF20-
-73.1	0.04	CF2ClCF2O-
- 78.6	a	$-oc_{\underline{F}_2}c_{\underline{F}}(c_{\underline{F}_3})o-$?
-79.0	b	$-oc_{\underline{F}_2}c_{\underline{F}}(c_{\underline{F}_3})o-$?
-80.3	b(broad)	?
-80.9	b	CF3CF2CF2O- ?
-81.8	b	-oc <u>F</u> 2cF2cF2c <u>F</u> 2o-
-82.1	b	cF ₃ cF ₂ c <u>F</u> ₂ o- ?
-85.1	ā	-OCF(C <u>F</u> 3)O-
-86.2	0.1	c <u>F</u> 3cF20-
-87.3	0.2	$cF_3c\underline{F}_2o-$, $-ocF_2ocF_2c\underline{F}_2oc\underline{F}_2cF_2cF_2ocF_2o-$
-89.0	4.8	-OCF ₂ OCF ₂ CF ₂ OCF ₂ O (major), -OCF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₂ O-
-89.5	0.1?	cf ₃ oc <u>f</u> 2cf ₂ o-
-89.9	a	?
-124.0	a	-OCF ₂ CF ₂ CF ₂ CF ₂ O-

a Peak is of extremely low intensity.

b Peak intensity cannot be determined due to poor resolution or low intensity.

TABLE 59 19 F NMR of XO[CF2CF20]nX (MLO 91-161)

ppm	Rel.Int.	Assignment			
-54.2	a	$CF_3OCF_2CF(CF_3) - ?$			
-54,8	0.9	CF3OCF2CF2O-			
-77.9	a	-00_{2}^{2} CF(0_{3}^{2})0- ?			
-78.9	a	$-0C\underline{F}_2CF(C\underline{F}_3)O-?$			
-80.6	b	CF ₃ CF ₂ CF ₂ 0-			
-81.5	b	CF ₃ CF ₂ CF ₂ O- ?			
-83.0	b	?			
-83.4	b	?			
-85.1	a	$-OCF(C\underline{F}_3)O-?$			
-86.1	0.2	$C\underline{F}_3CF_2O-$			
- 87.3	12.0	$-OC_{\underline{F}_2}C_{\underline{F}_2}O- (major), CF_3C_{\underline{F}_2}O-$			
-88.1	a	? (major), crac <u>r</u> 20-			
-89.4	0.6	CF ₃ OC <u>F₂</u> CF ₂ O-			
-136.6	a	?			

a Peak is of extremely low intensity

b Peak intensity cannot be determined due to the poor resolution or low intensity.

TABLE 60

SEPARATION AND CHARACTERIZATION DATA FOR SELECTED PERFLUOROALKYLETHER FLUIDS^a

Material	BP, °C	Viscosi 40°C	ty, cSt	VI	MW (Osm)
-(CF ₂ CF ₂ CF ₂ O) _n - Demnum S-20	As Received	23.8	5.48	179	2500
Fraction 1	86-142/0.001 mm Hg	9.24	2.44	77	1740
Fraction 2	142-175/0.001 mm Hg	14.3	3.48	123	2190
Fraction 3	175-225/0.001 mm Hg	24.7	5.48	168	2940
Residue	>225/0.001 mm Hg	80.6	15.5	205	5410
-(CF ₂ CF ₂ CF ₂ CF ₂ OCF ₂ O) _n - (MLO 91-105)	180-330/0.5 mm Hg	20.8	5.04	183	2500
Fraction 1	114-150/0.001 mm Hg	10.8	2.89	119	1960
Fraction 2	150-162/0.001 mm Hg	13.3	3.44	140	2190
Fraction 3	162-201/0.001 mm Hg	19.0	4.64	172	2760
Residue	>201/0.001 mm Hg	36.1	8.27	215	4030
-(CF ₂ CF ₂ CF ₂ CF ₂ O) _n - (MLO 91-126)	180-330/0.5 mm Hg	52.0	7.73	114	2600
Fraction 1	50-170/0.001 mm Hg	21.2	3.88	52	1800
Fraction 2	170-206/0.001 mm Hg	42.4	6.50	103	2570
Fraction 3	206-231/0.001 mm Hg	64.5	8.97	114	3180
Residue	>231/0.001 mm Hg	116.6	14.5	126	4210
-[CF ₂ CF ₂ OCF ₂ CF(CF ₂ CF ₃)OCF ₂ O] _n -					
(MLO 91-131)	180-330/0.5 mm Hg	22.6	5.00	155	2900
Fraction 1	52-164/0.001 mm Hg	11.3	2.82	90	2160
Fraction 2	164-190/0.001 mm Hg	18.1	4.11	131	2800
Fraction 3	190-212/0.001 mm Hg	25.3	5.39	155	3260
Residue	>212/0.001 mm Hg	44.0	8.58	177	4630
-[(CF ₂ CF(CF ₃)) ₂ OCF ₂ O] _n - (MLO 91-132)	180-330/0.5 mm Hg	28.4	6.00	165	3050
Fraction 1	130-163/0.001 mm Hg	15.3	3.50	106	2300
Fraction 2	163-189/0.001 mm Hg	24.8	5.22	148	3200
Residue	>189/0.001 mm Hg	51.6	9.76	178	4070
$-[(CF_2CF_2O)_4CF_2O]_n$ (MLO 91-157)	180-330/0.5 mm Hg	15.4	3.95	162	2870
Fraction 1	50-157/0.001 mm Hg	7.38	2.17	93	1970
Fraction 2	157-166/0.001 mm Hg	10.9	2.96	128	2570
Fraction 3	166-194/0.001 mm Hg	14.6	3.74	152	2870
Residue	>194/0.001 mm Hg	28.4	6.40	188	4350

SEPARATION AND CHARACTERIZATION DATA FOR SELECTED PERFLUOROALKYLETHER FLUIDS^a

TABLE 60 .(CONCLUDED)

Material	BP, °C		ty, cSt	VI	MW
		40°C	100°C		(Osm)
-[CF ₂ CF(CF ₂ OCF ₂ CF ₂ OCF ₂ CF ₂ OCF ₃)O] _n -					
(MLO 91-158)	180-330/0.5 mm Hg	67.1	8.71	102	3950
Fraction 1	70-161/0.001 mm Hg	25.6	4.21	35	2770
Fraction 2	161-189/0.001 mm Hg	45.2	6.34	84	3350
Fraction 3	189-220/0.001 mm Hg	81.8	9.73	96	4310
Residue	>220/0.001 mm Hg	169.5	18.0	117	6300
-(CF ₂ CF ₂ OCF ₂ O) _n - (MLO 91-160)	180-330/0.5 mm Hg	10.5	3.32	215	2640
Fraction 1	99-147/0.001 mm Hg	5.96	2.03	149	2030
Fraction 2	147-166/0.001 mm Hg	7.81	2.58	187	2470
Fraction 3	166-186/0.001 mm Hg	9.98	3.19	213	2700
Residue	>186/0.001 mm Hg	17.4	5.12	257	3850
-(CF ₂ CF ₂ O) _n - (MLO 91-161)	180-330/0.5 mm Hg	16.7	4.04	141	3000
Fraction 1	120-160/0.001 mm Hg	10.0	2.68	104	2400
Fraction 2	160-173/0.001 mm Hg	12.7	3.24	124	2800
Residue	>173/0.001 mm Hg	26.9	5.78	166	4250

a) The original material as received from WL/MLBT was redistilled at 0.001 mm Hg and two or three fractions plus a residue was obtained

TABLE 61 page 1 of 5
COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PERFLUOROALKYLETHER FLUIDS^a

Test No.	Fluid Type	g	Temp.	Volatiles mg/g	Coupon Weight Change, mg/cm ²
1	Krytox (MLO 71-6)	3.02	260	0.13	+0.1
28b	ditto	9.99	300(7	72h)38.2	+0.4
32	ditto	3.41	315	44.0	+0.5
105	Aflunox (MLO 88-298)	3.09	315	0.19	+0.1
93	ditto	3.03	325	5.5	+0.4
106	ditto	3.07	330	12.7	+0.7
2	Fomblin Z (Z25-P28)	3.17	260	57.4	-0.2
94	Fomblin Z (Z25-P151)	3.05	315	3.4	no coupon
98	ditto	3.06	315	5.5	no coupon ^r
3	Demnum S-20 (MLO 86-51)	3.74	260	1.0	+0.2
27 ^C	Demnum S-20 (MLO 88-177)	9.94	260(58h) 0.7	n.d.
29d	Aged Demnum S-20	9.93	300	1.4	+3.1
831	ditto	2.90	300	0.07	n.d.
41	Demnum S-20	3.05	315	16.8	+1.1
78	ditto	6.38	315	10.4	+1.2
79 ⁱ	Aged Demnum S-20	3.24	315	5.4	+0.7
82k	ditto	2.99	315	0.70	0.0
77	Demnum S-20	3.05	330	37.8	+1.0
31e	Aged Demnum S-20	3.29	330	13.3	+0.8

TABLE 61 (continued)

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COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PERFLUOROALKYLETHER
FLUIDS^a

Test No.	Fluid Type	g :	Temp.	Volatiles mg/g	Coupon Weight Change, mg/cm ²
80j	Aged Demnum S-20	3.10	330	5.3	+1.2
84m	ditto	2.90	345	73.7	+0.3
53	Demnum S-20	3.03	345	2.6	no coupon
99	Demnum S-100	3.22	315	8.0	+0.54
100r	ditto	3.20	315	4.7	$(Si_3N_4) + 0.48$
102 ^t	Aged Demnum S-100	3.09	315	0.65	+0.18
101 ^s	ditto	3.16	330	6.1	+2.4
104	Demnum S-100	3.13	330	22.5	+1.8
4	Brayco 814Z (MLO 78-80)	3.32	260	4.8	+0.2
35	ditto	3.36	275	62.2	+0.4
38	Fluor. Krytox (MLO 91-21)	3.08	315	0.03	+0.1
89	ditto	3.05	325	0.3	+0.06
870	ditto	3.03	325(40	h) 3.8	-0.2
55	ditto	3.02	330	2.0	+0.2
46	ditto	3.08	345	27.4	-0.1
52	ditto	3.03	345	0.99	no coupon
57	ditto	3.01	360	4.8	no coupon
56 ^h	ditto	3.05	360 (N ₂) 0.69	0.0
36	MLO 91-87f	3.20	300	0.06	+0.1
39	ditto	3.04	315	0.59	+0.1

TABLE 61 (continued)

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COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PERFLUOROALKYLETHER

FLUIDS^a

Test	Fluid Type	g	Temp.	Volatiles mg/g	Coupon Weight Change, mg/cm ²
48	MLO 91-87	3.05	330	55.6	0.0
37	MLO 91-889	3.20	300	0.09	+0.1
40	ditto	3.14	315	2.2	+0.1
49	ditto	3.04	330	121.6	-0.2
59 -	[(CF ₂)50CF ₂ 0] _n - (MLO 91-10	06)3.01	300	0.53	-0.06
96	ditto	3.05	310	0.10	0.0
50	ditto	3.15	315	11.9	+0.1
42	ditto	3.18	330	55.0	+0.1
60 -	[(CF ₂) ₄ 0] _n - (MLO 91-126)	3.03	315	0.99	-0.06
86 ⁿ	Aged MLO 91-126	2.82	315	0.71	0.0
91P	ditto	2.87	315	1.6	+0.4
43	MLO 91-126	3.06	330	5.1	+0.06
47	ditto	3.04	345	73.8	+0.4
58-[(CF ₂ CF ₂ O) ₄ CF ₂ O] _n - (MLO91-15	57)3.02	280	0.53	0.0
54	ditto	3.02	290	3.7	-0.1
44	ditto	3.02	300	25.7	+0.4
45	[CF ₂ CF ₂ O] _n (MLO 91-161)	3.04	300	0.26	0.0
929	ditto	2.76	300 (4	18h)28.9	+0.8
61	ditto	3.04	310	0.62	0.0
51	ditto	3.01	315	18.06	0.0

TABLE 61 (continued)

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COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PERFLUOROALKYLETHER
FLUIDSa

Test	Fluid Type	g	Temp. °C	Volatiles mg/g	Coupon Weight Change, mg/cm ²
62-[0	F ₂ CF(CF ₃)CF ₂ CF(CF ₃)OCF ₂ O] _r	 n-3.03	300	0.56	+0.06
107 ^v	Aged MLO 91-132	2.80	310	59.6	+0.06
64	MLO 91-132	3.07	315	5.0	+0.2
66	ditto	3.04	325	33.6	+0.2
63 -[CF ₂ CF[(OCF ₂ CF ₂) ₂ OCF ₃]0] _n - (MLO 91-158)	3.02	300	0.13	-0.06
85	ditto	3.01	305	0.30	0.0
95	ditto	3.01	310	0.03	0.0
65	ditto	3.02	315	9.0	+0.1
67	ditto	3.05	325	175.0	-1.4
68 -[$(CF_2)_4OCF_2O]_n$ - (MLO 91-105	3.06	300	0.13	0.0
81	ditto	3.05	305	0.07	-0.06
88	ditto	3.02	310	0.30	+0.06
70	ditto	3.06	315	9.1	-0.06
76	ditto	3.04	330	31.5	+0.06
72 -(CF ₂ CF ₂ OCF ₂ O) _n - (MLO 91-160)3.04	275	0.06	0.0
90	ditto	3.01	285	0.03	0.0
75	ditto	3.06	290	2.2	-0.06
69	ditto	3.04	300	118.4	-0.5
71 -[CF ₂ CF(CF ₃)0] _n - (MLO 91-127)3.01	315	0.30	0.0

TABLE 61 (concluded)

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COMPARISON OF THERMAL OXIDATIVE BEHAVIOR OF VARIOUS PERFLUOROALKYLETHER
FLUIDS^a

Test No.	Fluid Type	g	Temp.	Volatiles mg/g	Coupon Weight Change, mg/cm ²
74 -[CF	C2CF(CF3)0]n- (MLO 91-	-127)3.05	325	1.6	+0.06
73 d	litto	3.06	330	18.8	-0.06

- (a) All the tests were performed in pure oxygen at the denoted temperature in the presence of an M-50 coupon over a 24 h period using the modified, scaled-down, sealed version of the AFML Micro-O-C-Test arrangement.
- (b) Test conducted for 72 h.
- (c) Test conducted for 68 h.
- (d) Fluid and coupon from Test 27 used for this test.
- (e) Fluid from Test 29 used for this test.
- (f) MLO 91-87, $CF_3O[CF_2CF_2O]_m[CF_2CF(CF_3)O]_n[CF_2CF_2O]_pCF_3$
- (g) MLO-91-88, $CF_2^{2}O[CF_2^{2}CF(CF_3^{2})O]_n^{-}[CF_2CF_2O]_mCF_3$

CFO[CF₂CF(CF₃)0]_q[CF₂CF₂0]_pCF₃

CF20[CF2CF(CF3)0]r[CF2CF20]sCF3

- (h) Test conducted in nitrogen.
- (i) Fluid from Test 78 used for this test.
- (j) Fluid from Test 79 used for this test.
- (k) Fluid from Test 80 used for this test.
- (1) Fluid from Test 82 used for this test.
- (m) Fluid from Test 85 used for this test.
- (n) Fluid from Test 60 used for this test.
- (o) This test was conducted over a 40 h period.
- (p) Fluid from Test 43 used for this test.
- (q) This test was conducted over a 48 h period.
- (r) Si3N4 ball used in this test.
- (s) Fluid from Test 99 used for this test.
- (t) Fluid from Test 100 used for this test.
- (u) Fluid from Test 101 used for this test.
- (v) Fluid from Test 64 used for this test.

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